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L1
                 SEL RN
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L2
L3
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              0 S L2 AND NCNC2-SC4/ES
L4
              6 S L2 AND P/ELS
L5
             86 S L2 AND SQL/FA
L6
             17 S L6 AND 11/SQL
L7
             26 S L6 AND 12/SQL
\Gamma8
              4 S L8 AND PEPTIDE NUCLEIC ACID AND THIENO AND IMIDAZOL AND HEXAH
Ь9
              1 S L9 AND G G T A T G G G A T A T
L10
                E FS
                E GGTATGGGATAT/SQEN
              3 S E3
L11
                 E TATTCCGTCAT/SQEN
            129 S E3
L12
              4 S L12 AND THIENO AND IMIDAZOL?
L13
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L22
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L29
            127 S L28 AND OXOPENTYL AMINO HEXYL
L30
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L41
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L42
                                                                      Jan Delaval
L43
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L45
             14 S L12 AND ?PHOSPHINYL?/CNS
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                                                                  jan.delaval@uspto.gov
L47
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L48
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2 S L48 NOT ?THIENO?/CNS

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L77
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T80
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L81
           9 S L79,L80
L82
           3 S L78 NOT L81
=> fil reg .
FILE 'REGISTRY' ENTERED AT 08:50:05 ON 13 MAR 2003
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STRUCTURE FILE UPDATES: 12 MAR 2003 HIGHEST RN 498527-50-7 DICTIONARY FILE UPDATES: 12 MAR 2003 HIGHEST RN 498527-50-7

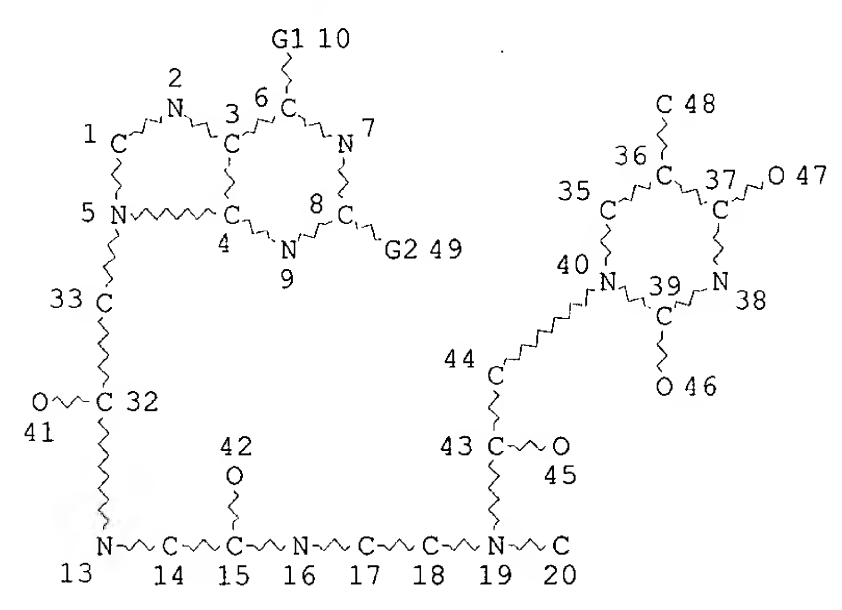
TSCA INFORMATION NOW CURRENT THROUGH MAY 20, 2002

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Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

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VAR G1=O/N
VAR G2=H/N
NODE ATTRIBUTES:
CONNECT IS M1 RC AT 13
CONNECT IS M1 RC AT 20
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 35

STEREO ATTRIBUTES: NONE
L52 157 SEA FILE=REGISTRY CSS FUL L50

100.0% PROCESSED 342 ITERATIONS 157 ANSWERS SEARCH TIME: 00.00.01

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FILE COVERS 1907 - 13 Mar 2003 VOL 138 ISS 11 FILE LAST UPDATED: 12 Mar 2003 (20030312/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

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L69 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2003 ACS
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- AN 2001:780930 HCAPLUS
- DN 135:331678
- TI Methods for preparing phosphorylated peptide nucleic acids carrying one or more marker, crosslinking, intracellular uptake, or binding affinity groups
- IN Uhlmann, Eugen; Breipohl, Gerhard; Will, David William
- PA Aventis Pharma Deutschland G.m.b.H., Germany
- SO PCT Int. Appl., 96 pp.
 - CODEN: PIXXD2
- DT Patent
- LA German
- IC ICM C07H021-00
- CC 34-3 (Amino Acids, Peptides, and Proteins)
 Section cross-reference(s): 33

FAN.CNT 1

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PRAI DE 2000-10019136 A
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    WO 2001-EP4027
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AB The invention relates to PNA derivs. which carry a phosphoryl radical on the N terminus of the PNA backbone, for example a phosphate or a substituted phosphoryl radical, substituted phosphoryl derives optionally carrying one or more marker groups or groups for crosslinking or groups which favor intracellular take-up or groups which increase the binding affinity of the PNA deriv. to nucleic acids. The invention also relates to a method for producing the aforementioned PNA derivs. and to their use

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as medicaments and diagnostic agents. Thus, several PNA chains were prepd.using solid phase peptide synthesis techniques, in which the C-terminal was capped by NH(CH2)60H and the N-terminal H2N- group was replaced by HO-, and functionalized to H2O3PO- or ROP(O)(OH)O- (R = biotin or fluorescein tag group or alkyl cap). Hybridization tests with complementary DNA or RNA showed increased binding, compared to a normal PNA chain N-capped with H3CC(0) - and C-capped with NH(CH2)60H. In vitro cellular uptake studies were done with fluorescein-tagged PNA (no data). In vitro cell proliferation studies were done with a H3C(CH2)15OP(O)(OH)capped PNA using human pre-B leukemia cells or A549-tumor cells (no data). PNA deriv prepn antiviral antimicrobial antitumor diagnostic hybridization Diagnosis (agents; prepn. of PNA derivs. as therapeutic or diagnostic agents) Solid phase synthesis (peptide; prepn. of PNA derivs. as therapeutic or diagnostic agents) Antimicrobial agents Antitumor agents Antiviral agents Biosensors Nucleic acid hybridization (prepn. of PNA derivs. as therapeutic or diagnostic agents) Peptide nucleic acids RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 368944-36-9P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 368944-38-1P 368944-39-2P 368944-40-5P 368944-41-6P 368944-42-7P 368944-43-8P 368944-44-9P 368944-45-0P RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 368506-25-6P **368944-35-8P** 368944-37-0P RL: RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 367255-38-7P 367255-39-8P 367985-52-2P 367985-53-3P 367985-54-4P 367985-55-5P **368506-26-7P 368506-27-8P 368506-28-9P 368506-29-0P** 368506-30-3P 368506-31-4P 368944-46-1P RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 110616-00-7 116364-61-5 147178-75-4 159845-57-5 169025-57-4, GenBank AR029142 181988-02-3 181988-09-0 185831-42-9 186070-79-1, GenBank A42375 186071-78-3 186108-31-6, 3: PN: WO0004034 SEQID: 3 unclaimed DNA 186123-93-3, GenBank A44395 186162-52-7 186162-55-0, GenBank A42368 189356-60-3 195184-07-7, GenBank A42342 195184-11-3, GenBank A42347 195184-12-4 195184-14-6, GenBank A42351 195184-15-7, GenBank A42352 195184-16-8, GenBank A44386 195184-17-9, GenBank A42354 195184-18-0, GenBank A42355 195184-19-1, GenBank A42356 195184-20-4, GenBank A42357 195184-21-5, GenBank A42358 195184-22-6, GenBank A42359 195184-23-7, GenBank A42361 195184-24-8, GenBank A42362 195184-25-9, GenBank A42363 195184-26-0, GenBank A47186 195184-27-1 195184-28-2, GenBank A47179 197103-72-3 197831-18-8 246223-25-6 257601-47-1, GenBank AX283184 325605-36-5, GenBank AX283169 325605-37-6, GenBank AX283174 325605-38-7 325605-39-8 325605-40-1 325605-41-2

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     RL: PRP (Properties)
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                            COPYRIGHT 2003 ACS
L69 ANSWER 2 OF 3 HCAPLUS
     2001:780897 HCAPLUS
ΑN
    135:331677
DN
    Methods for preparing phosphorylated peptide nucleic acids carrying one or
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     more marker, crosslinking, intracellular uptake, or binding affinity
     groups
    Uhlmann, Eugen; Breipohl, Gerhard; Will, David
ΙN
     William
    Aventis Pharma Deutschland G.m.b.H., Germany
PA
     PCT Int. Appl., 93 pp.
SO
     CODEN: PIXXD2
DT
    Patent
    German
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     34-3 (Amino Acids, Peptides, and Proteins)
CC
     Section cross-reference(s): 6, 33, 63
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     WO 2001-EP4030
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OS
     The invention relates to PNA derivs. that carry one or more phosphoryl
AΒ
     groups at the C terminus or at the C and N terminus of the PNA backbone,
     said phosphoryl groups optionally carrying one or more marker groups, or
     groups for crosslinking, or groups that promote the intracellular uptake,
     or groups that improve the binding affinity of the PNA deriv. to nucleic
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acids. The invention further relates to a method for producing the above

Thus, title compd. CH3(CH2)15OP(O)(OH)-T(oeq)[ATTCCGTCAT](CH2)6NHP(O)(OH)O-

PNA derivs. and to the use thereof as a medicament or diagnostic agent.

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CH2CH(CH2OH)(CH2)4NHC(S)NH-fluorescein (I) [T(oeg) = O(CH2)2N(C(O)CH2-Base)CH2C(O)-; remainder of chain = normal peptide nucleic acid backbone] was prepd. using solid-phase peptide synthesis techniques. Hybridization tests of I with complementary DNA and RNA showed better complexation with DNA than with RNA, though both were stronger than with PNA Ac-NH-TATTCCGTCAT-(CH2)6NH2 ref. In vitro cell proliferation studies using I and human pre-B leukemia cells showed stronger inhibition than a known phosphorothioate oligonucleotide (no data). PNA deriv prepn antiviral antimicrobial antitumor diagnostic hybridization Diagnosis (agents; prepn. of PNA derivs. as therapeutic or diagnostic agents) Solid phase synthesis (peptide; prepn. of PNA derivs. as therapeutic or diagnostic agents) Antimicrobial agents Antitumor agents Antiviral agents Biosensors Nucleic acid hybridization (prepn. of PNA derivs. as therapeutic or diagnostic agents) Peptide nucleic acids RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 368505-39-9P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 367985-20-4P 367985-21-5P 367985-22-6P 367985-23-7P RL: PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 367985-17-9P 367985-19-1P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (prepn. of PNA derivs. as therapeutic or diagnostic agents) **367985-18-0P** 368505-37-7P **368505-38-8P** 368505-40-2P RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 110616-00-7 116364-61-5 147178-75-4 159845-57-5 169025-57-4, GenBank AR029142 181988-02-3 181988-09-0 186070-79-1, GenBank A42375 186123-93-3, GenBank A44395 186162-52-7 186162-55-0, GenBank A42368 189356-60-3 195184-07-7, GenBank A42342 195184-11-3, GenBank A42347 195184-12-4 195184-14-6, GenBank A42351 195184-15-7, GenBank A42352 195184-16-8, GenBank A44386 195184-17-9, GenBank A42354 195184-18-0, GenBank A42355 195184-19-1, GenBank A42356 195184-20-4, GenBank A42357 195184-21-5, GenBank A42358 195184-22-6, GenBank A42359 195184-23-7, GenBank A42361 195184-24-8, GenBank A42362 195184-25-9, GenBank A42363 195184-26-0, GenBank A47186 195184-27-1 195184-28-2, GenBank A47179 197831-18-8 246223-25-6 257601-47-1, GenBank AX283184 325605-36-5, GenBank AX283169 325605-37-6, GenBank AX283174 325605-38-7 325605-39-8 325605-40-1 325605-41-2 325605-42-3 325605-43-4 325605-44-5 325605-45-6 325605-46-7 325605-47-8 325605-48-9 325605-49-0 325605-50-3 325605-51-4 325605-52-5 RL: PRP (Properties) (unclaimed nucleotide sequence; methods for prepg. phosphorylated peptide nucleic acids carrying one or more marker, crosslinking,

intracellular uptake, or binding affinity groups)

- L69 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2003 ACS 2001:342363 HCAPLUS ΑN 135:122729 DN Synthesis of peptide nucleic acid oligomers carrying 5-methylcytosine ŢΙ derivatives by post synthetic substitution Ferrer, Elisenda; Eritja, Ramon ΑU European Molecular Biology Laboratory, Heidelberg, D-69117, Germany ÇŞ Letters in Peptide Science (2001), Volume Date 2000, 7(4), 195-206 SO CODEN: LPSCEM; ISSN: 0929-5666 Kluwer Academic Publishers PBJournal DT English LA34-3 (Amino Acids, Peptides, and Proteins) CC Section cross-reference(s): 33 CASREACT 135:122729 OS The prepn. of the thymine peptide nucleic acid (PNA) monomer carrying a AΒ 2-nitrophenyl group in position 4 is described. This monomer is incorporated into PNA oligomers and reacted with amines to yield PNA oligomers carrying 5-methylcytosine derivs. During the deprotection-modification step two side reactions were detected: degrdn. of PNA oligomer from the N-terminal residue and modification of N4-tert-butylbenzoyl cytosine residue. Protection of the N-terminal position and the use of N4-acetyl group for the protection of cytosine eliminate these side reactions. peptide nucleic acid methylcytosine prepn reaction amine STAmines, reactions IT RL: RCT (Reactant); RACT (Reactant or reagent) (synthesis of peptide nucleic acid oligomers carrying 5-methylcytosine derivs.) Peptide nucleic acids IT RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (synthesis of peptide nucleic acid oligomers carrying 5-methylcytosine derivs.) Substitution reaction IT (synthesis of peptide nucleic acid oligomers carrying 5-methylcytosine derivs. by post synthetic substitution) 5036-48-6, 1-(3-Aminopropyl)imidazole 5292-43-3, tert-Butyl bromoacetate IT14631-20-0, n4-Acetylcytosine 244764-39-4 244764-42-9 244764-45-2 244764-46-3 244764-47-4 272788-86-0 RL: RCT (Reactant); RACT (Reactant or reagent) (synthesis of peptide nucleic acid oligomers carrying 5-methylcytosine derivs.) 350608-24-1P 350608-25-2P 350608-28-5P 350608-29-6P 350608-33**-**2P IT350608-34-3P 350608-35-4P 350608-36-5P 350728-22-2P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (synthesis of peptide nucleic acid oligomers carrying 5-methylcytosine derivs.) 350608-27-4P 350608-30-9P 350608-32-1P **350608-37-6P** IT**350608-38-7P** 350608-39-8P 350608-40-1P 350608-41-2P 350608-42-3P 350608-43-4P 350608-44-5P 350608-45-6P 350608-46-7P 350608-47-8P 350608-48-9P 350728-23-3P 350728-24-4P RL: SPN (Synthetic preparation); PREP (Preparation)
- RE.CNT 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD RE
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derivs.)

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(synthesis of peptide nucleic acid oligomers carrying 5-methylcytosine

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- (19) Will, D; Tetrahedron 1995, V51, P12069 HCAPLUS
- (20) Xu, Y; J Org Chem 1992, V57, P3839 HCAPLUS

=> sel hit rn 169 E565 THROUGH E575 ASSIGNED

=> fil reg

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STRUCTURE FILE UPDATES: 12 MAR 2003 HIGHEST RN 498527-50-7 DICTIONARY FILE UPDATES: 12 MAR 2003 HIGHEST RN 498527-50-7

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Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details: http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf

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CN
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Source
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Not Given | WO2001079249
        lunclaimed
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modified base t-1
                                        modified thymidine
                                        3'-deoxy
modified base t-11
                                        3'-substituted
modified base t-11
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    2 a 3 c 1 g 5 t
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                ----- location ----- description
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                                        5'-ester
modified base t-1
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modified base t-1
modified base t-11
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modified base
               t-11
SEQ
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L83
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CN
    oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphos
    phinyl]oxy]]T-G-A-A-G-G-A-A-G-A-G-G)-(6-hydroxyhexyl)NH (9CI) (CA INDEX
    NAME)
    NUCLEIC ACID SEQUENCE
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 type ----- location ----- description
modified base t-1
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                                        modified thymidine
modified base t-1
                                        3'-deoxy
modified base g-12
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modified base g-12
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CI

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L83 ANSWER 5 OF 11 REGISTRY COPYRIGHT 2003 ACS
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NTE modified
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                                        5'-ester
modified base g-1 modified base t-12
                                       modified guanosine
                                       3'-deoxy
modified base t-12
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SEQ
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L83 ANSWER 6 OF 11 REGISTRY COPYRIGHT 2003 ACS
    368506-27-8 REGISTRY
RN
    Peptide nucleic acid, ([5'-deamino-5'-[[[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-
CN
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FS
SQL 12
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NA
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type ----- location ----- description
modified base g-1
                                        5'-ester
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modified base c-12
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SEQ 1 gctgatgtag tc
**RELATED SEQUENCES AVAILABLE WITH SEQLINK**
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LC STN Files: CA, CAPLUS, TOXCENTER, USPATFULL 1 REFERENCES IN FILE CA (1962 TO DATE) 1 REFERENCES IN FILE CAPLUS (1962 TO DATE) REFERENCE 1: 135:331678 L83 ANSWER 7 OF 11 REGISTRY COPYRIGHT 2003 ACS 368506-26-7 REGISTRY RN Peptide nucleic acid, ([[[6-[[5-[(3aS,4S,6aR)-hexahydro-2-oxo-1H-CN thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphosphinyl]-A-C-T-G-A-T-G-T-A-G-T-C)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME) NUCLEIC ACID SEQUENCE FS 12 SQL 3 a 2 c 3 g 4 t NA NTE modified type ----- location ----- description 5'-substituted modified base a-1 modified base c-12 3'-deoxy modified base c-12 3'-substituted SEQ 1 actgatgtag tc **RELATED SEQUENCES AVAILABLE WITH SEQLINK** Unspecified MF CIMAN SR CA LCSTN Files: CA, CAPLUS, TOXCENTER, USPATFULL 1 REFERENCES IN FILE CA (1962 TO DATE) 1 REFERENCES IN FILE CAPLUS (1962 TO DATE) REFERENCE 1: 135:331678 ANSWER 8 OF 11 REGISTRY COPYRIGHT 2003 ACS L83 **368505-38-8** REGISTRY RN Peptide nucleic acid, ([5'-deamino-5'-[[[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-CN oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphos phinyl]oxy]]T-A-T-T-C-C-G-T-C-A-T)-[2-(phosphonooxy)ethyl]NH (9CI) (CA INDEX NAME) NUCLEIC ACID SEQUENCE FS SQL 2 a 3 c 1 g 5 t NΑ NTE modified type ----- location ----- description modified base t-1 modified base t-1 modified base t-11 5'-ester modified thymidine 3'-deoxy modified base t-11 SEQ 1 tattccgtca t **RELATED SEQUENCES AVAILABLE WITH SEQLINK** MFUnspecified CIMAN SR CA LCSTN Files: CA, CAPLUS, TOXCENTER, USPATFULL 1 REFERENCES IN FILE CA (1962 TO DATE) 1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

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L83 ANSWER 9 OF 11 REGISTRY COPYRIGHT 2003 ACS
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type ----- location ----- description
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SEQ
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REFERENCE 1: 135:331677
    ANSWER 10 OF 11 REGISTRY COPYRIGHT 2003 ACS
L83
RN
    350608-38-7 REGISTRY
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CN
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    NUCLEIC ACID SEQUENCE
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SQL
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NA
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type ----- location ----- description
modified base g-1
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modified base c-4
                                        m5c
modified base c-4 modified base c-4
                                        3'-deoxy
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PAGE 1-A

PAGE 2-A

PAGE 2-B

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1 REFERENCES IN FILE CAPLUS (1962 TO DATE)

REFERENCE 1: 135:122729

L83 ANSWER 11 OF 11 REGISTRY COPYRIGHT 2003 ACS

RN **350608-37-6** REGISTRY

CN Peptide nucleic acid, (acetyl-G-T-A-m5C)-(6-hydroxyhexyl)NH (9CI) (CA

INDEX NAME)

FS NUCLEIC ACID SEQUENCE

SQL 4

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NTE modified

type	location	description
modified base modified base modified base modified base	g-1 c-4 c-4	5'-ac m5c 3'-deoxy 3'-substituted

SEQ 1 gtac

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LC STN Files: CA, CAPLUS

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 CH_2
 CH_2

PAGE 2-B

--- CH2-NHAC

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REFERENCE 1: 135:122729

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This file contains CAS Registry Numbers for easy and accurate substance identification.

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L84 ANSWER 1 OF 12 HCAPLUS COPYRIGHT 2003 ACS

AN 2001:780930 HCAPLUS

DN 135:331678

- TI Methods for preparing phosphorylated peptide nucleic acids carrying one or more marker, crosslinking, intracellular uptake, or binding affinity groups
- IN Uhlmann, Eugen; Breipohl, Gerhard; Will, David William
- PA Aventis Pharma Deutschland G.m.b.H., Germany
- SO PCT Int. Appl., 96 pp. CODEN: PIXXD2

Relateduences

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Patent
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       German
   LA
       ICM C07H021-00
   ΙÇ
       34-3 (Amino Acids, Peptides, and Proteins)
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                               DATE
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                         W
       The invention relates to PNA derivs. which carry a phosphoryl
       radical on the N terminus of the PNA backbone, for example a
       phosphate or a substituted phosphoryl radical, substituted phosphoryl
GALGO K
       derives optionally carrying one or more marker groups or groups for
       crosslinking or groups which favor intracellular take-up or groups which
       increase the binding affinity of the PNA deriv. to nucleic
       acids. The invention also relates to a method for producing the
       aforementioned PNA derivs. and to their use as medicaments and
       diagnostic agents. Thus, several PNA chains were prepd.using
       solid phase peptide synthesis techniques, in which the C-terminal was
       capped by (NH(CH2) 60H and the N-terminal H2N- group was replaced by HO-,
       and functionalized to H2O3PO- or ROP(O)(OH)O- (R = biotin or fluorescein
       tag group or alkyl cap). Hybridization tests with complementary DNA or
       RNA showed increased binding, compared to a normal PNA chain
       N-capped with H3CC(0) - and C-capped with NH(CH2)60H. In vitro cellular
       uptake studies were done with fluorescein-tagged PNA (no data).
       In vitro cell proliferation studies were done with a H3C(CH2)15OP(O)(OH)-
       capped PNA using human pre-B leukemia cells or A549-tumor cells
       (no data).
  ST
       PNA deriv prepn antiviral antimicrobial antitumor diagnostic
       hybridization
       Diagnosis
   IT
           (agents; prepn. of PNA derivs. as therapeutic or diagnostic
          agents)
       Solid phase synthesis
  ΙT
           (peptide; prepn. of PNA derivs. as therapeutic or diagnostic
          agents)
       Antimicrobial agents
   IT
       Antitumor agents
       Antiviral agents
       Biosensors
       Nucleic acid hybridization
           (prepn. of PNA derivs. as therapeutic or diagnostic agents)
       Peptide nucleic acids
  IT
       RL: BAC (Biological activity or effector, except adverse); BSU (Biological
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RN

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study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
(Reactant or reagent); USES (Uses)
   (prepn. of PNA derivs. as therapeutic or diagnostic agents)
368944-36-9P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
(Reactant or reagent); USES (Uses)
   (prepn. of PNA derivs. as therapeutic or diagnostic agents)
368944-38-1P 368944-39-2P 368944-40-5P
368944-41-6P 368944-42-7P 368944-43-8P
368944-44-9P 368944-45-0P
RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
   (prepn. of PNA derivs. as therapeutic or diagnostic agents)
368506-25-6P 368944-35-8P 368944-37-0P
RL: RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use);
BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent);
USES (Uses)
   (prepn. of PNA derivs. as therapeutic or diagnostic agents)
367255-38-7P 367255-39-8P 367985-52-2P 367985-53-3P 367985-54-4P
367985-55-5P 368506-26-7P 368506-27-8P
368506-28-9P 368506-29-0P 368506-30-3P 368506-31-4P
368944-46-1P
RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
study); PREP (Preparation); USES (Uses)
   (prepn. of PNA derivs. as therapeutic or diagnostic agents)
110616-00-7 116364-61-5 147178-75-4 159845-57-5
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186070-79-1, GenBank A42375 186071-78-3 186108-31-6, 3: PN: WO0004034
SEQID: 3 unclaimed DNA 186123-93-3, GenBank A44395 186162-52-7
                             189356-60-3 195184-07-7, GenBank A42342
186162-55-0, GenBank A42368
                             195184-12-4 195184-14-6, GenBank A42351
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                             195184-16-8, GenBank A44386 195184-17-9,
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325605-46-7 325605-47-8 325605-48-9 325605-49-0 325605-50-3
325605-51-4 325605-52-5 368952-79-8 368952-80-1
                                                      368952-81-2
368952-82-3 368952-83-4 368952-84-5 368952-85-6
RL: PRP (Properties)
   (unclaimed nucleotide sequence; methods for prepg. phosphorylated
  peptide nucleic acids carrying one or more
  marker, crosslinking, intracellular uptake, or binding affinity groups)
81742-60-1 143189-17-7
RL: PRP (Properties)
   (unclaimed sequence; methods for prepg. phosphorylated peptide
  nucleic acids carrying one or more marker,
   crosslinking, intracellular uptake, or binding affinity groups)
368944-36-9P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
(Reactant or reagent); USES (Uses)
   (prepn. of PNA derivs. as therapeutic or diagnostic agents)
368944-36-9 HCAPLUS
Peptide nucleic acid, ([5'-deamino-5'-[[(hexadecyloxy)hydroxyphosphinyl]ox
```

v]]T-A-T-T-C-C-G-T-C-A-T)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME)

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*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
    368944-38-1P 368944-39-2P 368944-40-5P
     368944-41-6P 368944-42-7P 368944-43-8P
     368944-44-9P 368944-45-0P
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     368944-38-1 HCAPLUS
RN
     DNA, d(A-T-G-A-C-G-G-A-A-T-A), complex with peptide nucleic acid
CN
     ([5'-deamino-5'-(phosphonooxy)]T-A-T-T-C-C-G-T-C-A-T)-(6-hydroxyhexyl)NH
     (1:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
    368944-39-2 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-(phosphonooxy)]T-A-T-T-C-C-G-T-C-A-
CN
     T)-(6-hydroxyhexyl)NH, complex with RNA (A-U-G-A-C-G-G-A-A-U-A) (1:1)
     (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
RN
     368944-40-5 HCAPLUS
     DNA, d(A-T-G-A-C-G-G-A-A-T-A), complex with peptide nucleic acid
CN
     ([5'-deamino-5'-[[(hexadecyloxy)hydroxyphosphinyl]oxy]]T-A-T-T-C-C-G-T-C-A-
     T)-(6-hydroxyhexyl)NH (1:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
RN
     368944-41-6 HCAPLUS
     Peptide nucleic acid, ([5'-deamino-5'-[[(hexadecyloxy)hydroxyphosphinyl]ox
CN
     y]]T-A-T-T-C-C-G-T-C-A-T)-(6-hydroxyhexyl)NH, complex with RNA
     (A-U-G-A-C-G-G-A-A-U-A) (1:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368944-42-7 HCAPLUS
RN
     DNA, d(A-T-G-A-C-G-G-A-A-T-A), complex with peptide nucleic acid
CN
     ([5'-deamino-5'-[[[[6-[[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-
     1(3H),9'-[9H]xanthen]-5-yl)amino]thioxomethyl]amino]hexyl]oxy]hydroxyphosp
     hinvl[oxy]]T-A-T-T-C-C-G-T-C-A-T)-(6-hydroxyhexyl)NH (1:1) (9CI) (CA)
     INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368944-43-8 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-[[[[6-[[[(3',6'-dihydroxy-3-
CN
     oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5-
     yl)amino]thioxomethyl]amino]hexyl]oxy]hydroxyphosphinyl]oxy]]T-A-T-T-C-C-G-
     T-C-A-T)-(6-hydroxyhexyl)NH, complex with RNA (A-U-G-A-C-G-G-A-A-U-A)
     (1:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368944-44-9 HCAPLUS
RN
     DNA, d(A-T-G-A-C-G-G-A-A-T-A), complex with peptide nucleic acid
CN
     ([5'-deamino-5'-[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-oxo-1H-thieno[3, 4-
     d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphosphinyl]oxy]]T-A-T-
     T-C-C-G-T-C-A-T)-(6-hydroxyhexyl)NH (1:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368944-45-0 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-[[[6-[[5-[(3aS,4S,6aR)-hexahydro-2-
CN
     oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphos
     phinyl]oxy]]T-A-T-T-C-C-G-T-C-A-T)-(6-hydroxyhexyl)NH, complex with RNA
     (A-U-G-A-C-G-G-A-A-U-A) (1:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368506-25-6P 368944-35-8P 368944-37-0P
\operatorname{IT}
     RL: RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use);
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BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent);
     USES (Uses)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     368506-25-6 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-(phosphonooxy)]T-A-T-T-C-C-G-T-C-A-
CN
     T)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
    368944-35-8 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-[[[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-
CN
     oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphos
     phinyl]oxy]]T-A-T-T-C-C-G-T-C-A-T)-(6-hydroxyhexyl)NH (9CI) (CA INDEX
     NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
    368944-37-0 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-[[[6-[[(3',6'-dihydroxy-3-
CN
     oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5-
     yl)amino]thioxomethyl]amino]hexyl]oxy]hydroxyphosphinyl]oxy]]T-A-T-T-C-C-G-
     T-C-A-T)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368506-26-7P 368506-27-8P 368506-28-9P
\operatorname{IT}
     368506-29-0P
     RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
     study); PREP (Preparation); USES (Uses)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     368506-26-7 HCAPLUS
RN
CN
     Peptide nucleic acid, ([[[6-[[5-[(3aS,4S,6aR)-hexahydro-2-oxo-1H-
     thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphosphinyl]-
     A-C-T-G-A-T-G-T-A-G-T-C)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368506-27-8 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-[[[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-
CN
     oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphos
     phinyl]oxy]]G-C-T-G-A-T-G-T-A-G-T-C)-(6-hydroxyhexyl)NH (9CI) (CA INDEX
     NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368506-28-9 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-[[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-
CN
     oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphos
     phinyl]oxy]]G-G-T-A-T-G-G-G-A-T-A-T)-(6-hydroxyhexyl)NH (9CI) (CA INDEX
     NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     368506-29-0 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-[[[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-
CN
     oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphos
     phinyl]oxy]]T-G-A-A-G-G-A-A-G-A-G-G)-(6-hydroxyhexyl)NH (9CI) (CA INDEX
     NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
     116364-61-5 368952-84-5
IT
     RL: PRP (Properties)
        (unclaimed nucleotide sequence; methods for prepg. phosphorylated
       peptide nucleic acids carrying one or more
        marker, crosslinking, intracellular uptake, or binding affinity groups)
     116364-61-5 HCAPLUS
RN
     DNA, d(T-A-T-T-C-C-G-T-C-A-T) (9CI) (CA INDEX NAME)
CN
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Absolute stereochemistry.

PAGE 1-A

PAGE 2-B

PAGE 3-B

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RN 368952-84-5 HCAPLUS
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CN DNA, d(G-G-T-A-T-G-G-G-A-T-A-T) (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

L84 ANSWER 2 OF 12 HCAPLUS COPYRIGHT 2003 ACS

AN 2001:780897 HCAPLUS

DN 135:331677

- TI Methods for preparing phosphorylated **peptide nucleic acids** carrying one or more marker, crosslinking, intracellular uptake, or binding affinity groups
- IN Uhlmann, Eugen; Breipohl, Gerhard; Will, David
 William
- PA Aventis Pharma Deutschland G.m.b.H., Germany

SO PCT Int. Appl., 93 pp.

CODEN: PIXXD2

DT Patent

LA German

IC ICM CO7H

CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 6, 33, 63

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ΡI	WO	2001	0792	16	A	2	2001	1025		M	20	01-E	P403	0	2001	0407		
	WO 2001079216			A	A3 20020228													
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			HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,
			LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,
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             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
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                       Al
                                            NO 2002-4959
     NO 2002004959
                             20021015
                                                              20021015
                             20000418
PRAI DE 2000-10019135 A
     WO 2001-EP4030
                             20010407
     MARPAT 135:331677
OS
     The invention relates to PNA derivs. that carry one or more
AΒ
     phosphoryl groups at the C terminus or at the C and N terminus of the
     PNA backbone, said phosphoryl groups optionally carrying one or
     more marker groups, or groups for crosslinking, or groups that promote the
     intracellular uptake, or groups that improve the binding affinity of the
     PNA deriv. to nucleic acids. The invention further relates to a
     method for producing the above PNA derivs. and to the use
     thereof as a medicament or diagnostic agent. Thus, title compd.
     CH3(CH2)15OP(O)(OH)-T(oeg)[ATTCCGTCAT](CH2)6NHP(O)(OH)O-
     CH2CH(CH2OH)(CH2)4NHC(S)NH-fluorescein (I) [T(oeg) = O(CH2)2N(C(O)CH2-CH2CH(CH2OH)(CH2)4NHC(S)NH-fluorescein (I) <math>[T(oeg) = O(CH2)2N(C(O)CH2-CH2)]
     Base) CH2C(O) -; remainder of chain = normal peptide
     nucleic acid backbone] was prepd. using solid-phase
     peptide synthesis techniques. Hybridization tests of I with complementary
     DNA and RNA showed better complexation with DNA than with RNA, though both
     were stronger than with PNA Ac-NH-TATTCCGTCAT-(CH2)6NH2 ref. In
     vitro cell proliferation studies using I and human pre-B leukemia cells
     showed stronger inhibition than a known phosphorothicate oligonucleotide
     (no data).
     PNA deriv prepn antiviral antimicrobial antitumor diagnostic
ST
     hybridization
     Diagnosis
ΙT
        (agents; prepn. of PNA derivs. as therapeutic or diagnostic
        agents)
     Solid phase synthesis
IT
        (peptide; prepn. of PNA derivs. as therapeutic or diagnostic
        agents)
     Antimicrobial agents
IT
     Antitumor agents
     Antiviral agents
     Biosensors
     Nucleic acid hybridization
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     Peptide nucleic acids
IT
     RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic
     preparation); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); RACT (Reactant or reagent); USES (Uses)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     368505-39-9P
IT
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
     (Reactant or reagent); USES (Uses)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     367985-20-4P 367985-21-5P 367985-22-6P
IT
     367985-23-7P
     RL: PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
         (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     367985-17-9P 367985-19-1P
IT
```

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RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
    367985-18-0P 368505-37-7P 368505-38-8P
\operatorname{IT}
    368505-40-2P
    RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological
    study); PREP (Preparation); USES (Uses)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
    110616-00-7 116364-61-5 147178-75-4 159845-57-5
IT
    169025-57-4, GenBank AR029142 181988-02-3 181988-09-0 186070-79-1,
    GenBank A42375 186071-78-3 186108-31-6, 3: PN: WO0004034 SEQID: 3
    unclaimed DNA 186123-93-3, GenBank A44395 186162-52-7 186162-55-0,
    GenBank A42368 189356-60-3 195184-07-7, GenBank A42342 195184-11-3,
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    325605-36-5, GenBank AX283169 325605-37-6, GenBank AX283174
    325605-38-7 325605-39-8 325605-40-1 325605-41-2 325605-42-3
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    325605-48-9 325605-49-0 325605-50-3 325605-51-4 325605-52-5
    RL: PRP (Properties)
        (unclaimed nucleotide sequence; methods for prepg. phosphorylated
       peptide nucleic acids carrying one or more
       marker, crosslinking, intracellular uptake, or binding affinity groups)
    368505-39-9P
IT
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT
     (Reactant or reagent); USES (Uses)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     368505-39-9 HCAPLUS
RN
    Peptide nucleic acid, ([5'-deamino-5'-[[(hexadecyloxy)hydroxyphosphinyl]ox
CN
    y] T-A-T-T-C-C-G-T-C-A-T) - [17-[(3', 6'-dihydroxy-3-oxospiro[isobenzofuran-
    1(3H),9'-[9H]xanthen]-5-yl)amino]-8-hydroxy-11-(hydroxymethyl)-8-oxido-17-
     thioxo-7,9-dioxa-16-aza-8-phosphaheptadec-1-yl]NH (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
    367985-20-4P 367985-21-5P 367985-22-6P
IT
    367985-23-7P
    RL: PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use);
     BIOL (Biological study); PREP (Preparation); USES (Uses)
        (prepn. of PNA derivs. as therapeutic or diagnostic agents)
     367985-20-4 HCAPLUS
RN
    DNA, d(A-T-G-A-C-G-G-A-A-T-A), complex with peptide nucleic acid
CN
     ([5'-deamino-5'-(phosphonooxy)]T-A-T-T-C-C-G-T-C-A-T)-[17-[(3',6'-
     dihydroxy-3-oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5-yl)amino]-8-
     hydroxy-11-(hydroxymethyl)-8-oxido-17-thioxo-7,9-dioxa-16-aza-8-
    phosphaheptadec-1-yl]NH (1:1) (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
    367985-21-5 HCAPLUS
RN
     Peptide nucleic acid, ([5'-deamino-5'-(phosphonooxy)]T-A-T-T-C-C-G-T-C-A-
ÇN
    T)-[17-[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5-
     yl)amino]-8-hydroxy-11-(hydroxymethyl)-8-oxido-17-thioxo-7,9-dioxa-16-aza-
     8-phosphaheptadec-1-yl]NH, complex with RNA (A-U-G-A-C-G-G-A-A-U-A) (1:1)
     (9CI) (CA INDEX NAME)
*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***
```

367985-22-6 HCAPLUS

RN

DNA, d(A-T-G-A-C-G-G-A-A-T-A), complex with peptide nucleic acid CN ([5'-deamino-5'-[[(hexadecyloxy)hydroxyphosphinyl]oxy]]T-A-T-T-C-C-G-T-C-A-T)-[17-[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5yl)amino]-8-hydroxy-11-(hydroxymethyl)-8-oxido-17-thioxo-7,9-dioxa-16-aza-8-phosphaheptadec-1-yl]NH (1:1) (9CI) (CA INDEX NAME) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 367985-23-7 HCAPLUS RNPeptide nucleic acid, ([5'-deamino-5'-[[(hexadecyloxy)hydroxyphosphinyl]ox CN y]]T-A-T-T-C-C-G-T-C-A-T)-[17-[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-1(3H), 9'-[9H] xanthen]-5-y1) amino]-8-hydroxy-11-(hydroxymethy1)-8-oxido-17thioxo-7,9-dioxa-16-aza-8-phosphaheptadec-1-yl]NH, complex with RNA (A-U-G-A-C-G-G-A-A-U-A) (1:1) (9CI) (CA INDEX NAME) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 367985-17-9P 367985-19-1P IT RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 367985-17-9 HCAPLUS RNPeptide nucleic acid, (acetyl-T-A-T-T-C-C-G-T-C-A-T)-[6-CN (phosphonooxy)hexyl]NH (9CI) (CA INDEX NAME) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 367985-19-1 HCAPLUS RN Peptide nucleic acid, ([5'-deamino-5'-(phosphonooxy)]T-A-T-T-C-C-G-T-C-A-CN T)-[17-[(3',6'-dihydroxy-3-oxospiro[isobenzofuran-1(3H),9'-[9H]xanthen]-5yl)amino]-8-hydroxy-11-(hydroxymethyl)-8-oxido-17-thioxo-7,9-dioxa-16-aza-8-phosphaheptadec-1-yl]NH (9CI) (CA INDEX NAME) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 367985-18-0P 368505-37-7P 368505-38-8P 368505-40-2P RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (prepn. of PNA derivs. as therapeutic or diagnostic agents) 367985-18-0 HCAPLUS RNCN Peptide nucleic acid, ([5'-[[[(6-aminohexyl)oxy]hydroxyphosphinyl]oxy]-5'deamino]T-A-T-T-C-C-G-T-C-A-T)-[6-(phosphonooxy)hexyl]NH (9CI) (CA INDEX NAME) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 368505-37-7 HCAPLUS RNPeptide nucleic acid, (acetyl-T-A-T-T-C-C-G-T-C-A-[3'-de(carboxymethyl)-3'-CN [2-(phosphonooxy)ethyl]]T) (9CI) (CA INDEX NAME) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 368505-38-8 HCAPLUS RNPeptide nucleic acid, ([5'-deamino-5'-[[[6-[[5-[(3aS, 4S, 6aR)-hexahydro-2-CN oxo-1H-thieno[3,4-d]imidazol-4-yl]-1-oxopentyl]amino]hexyl]oxy]hydroxyphos phinyl]oxy]]T-A-T-T-C-C-G-T-C-A-T)-[2-(phosphonooxy)ethyl]NH (9CI) (CA INDEX NAME) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 368505-40-2 HCAPLUS RNCN Peptide nucleic acid, ([5'-[(28-amino-1,21-dihydroxy-1,21-dioxido-2,5,8,11,14,17,20,22-octaoxa-1,21-diphosphaoctacos-1-y1)oxy]-5'-deamino]T-A-T-T-C-C-G-T-C-A-T)-[6-(phosphonooxy)hexyl]NH, complex with RNA (A-U-G-A-C-G-G-A-A-U-A) (1:1) (9CI) (CA INDEX NAME) *** STRUCTURE DIAGRAM IS NOT AVAILABLE *** 116364-61-5 IT

RL: PRP (Properties)

(unclaimed nucleotide sequence; methods for prepg. phosphorylated peptide nucleic acids carrying one or more

marker, crosslinking, intracellular uptake, or binding affinity groups)

RN 116364-61-5 HCAPLUS

CN DNA, d(T-A-T-T-C-C-G-T-C-A-T) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 2-B

PAGE 3-B

DE 19935302

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ANSWER 3 OF 12 HCAPLUS COPYRIGHT 2003 ACS
L84
ΑN
     2001:101001 HCAPLUS
DN
     134:183461
     Conjugates and methods for the production thereof for transporting
TI
     molecules across biological membranes
     Uhlmann, Eugen; Greiner, Beate; Unger, Eberhard; Gothe,
ΙN
     Gislinde; Schwerdel, Marc
     Aventis Pharma Deutschland Gmbh, Germany
PΑ
SO
     PCT Int. Appl., 84 pp.
     CODEN: PIXXD2
     Patent
DT
     German
\mathtt{L}\mathtt{A}
IC
     ICM A61K047-48
     ICS A61K049-00
     63-5 (Pharmaceuticals)
CC
     Section cross-reference(s): 1, 9
FAN.CNT 1
     PATENT NO. KIND DATE
                                           APPLICATION NO.
                                                            DATE
     WO 2001008707 A2 20010208
                                           WO 2000-EP6936 20000720
PΙ
     WO 2001008707 A3
                            20011108
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU,
             ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
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CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

A1 20010208

DE 1999-19935302 19990728

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BR 2000012757
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                    A
                           20020515
    EP 1204430
                      A2
                                          EP 2000-956220
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        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL
                           20030212
                                          JP 2001-513437
    JP 2003505517
                      T2
                                                           20000720
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    NO 2002000367
                                          NO 2002-367
                                                           20020123
PRAI DE 1999-19935302 A
                          19990728
                           20000720
    WO 2000-EP6936
    MARPAT 134:183461
OS
    The invention relates to conjugates, methods for their prodn., and to the
AΒ
    use of these conjugates for transporting low mol. wt. compds. and
    macromols. across biol. membranes, in particular, for transporting mols.
    into cells. The invention also relates to medicaments, diagnostic agents
    and test kits in which these conjugates are present or introduced.
    drug delivery conjugate oligonucleotide membrane transport
ST
    Diagnosis
IT
        (agents; conjugates for transporting mols. across biol. membranes)
    Drug delivery systems
IT
        (carriers; conjugates for transporting mols. across biol. membranes)
    Antitumor agents
IT
    Bacteria (Eubacteria)
    Biological transport
    Cell membrane
    Eukaryote (Eukaryotae)
    Mammal (Mammalia)
    Molecular weight distribution
    Neoplasm
    Prokaryote
    Test kits
    Yeast
       (conjugates for transporting mols. across biol. membranes)
ΙT
    Macromolecular compounds
    RL: BPR (Biological process); BSU (Biological study, unclassified); PEP
     (Physical, engineering or chemical process); THU (Therapeutic use); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (conjugates; conjugates for transporting mols. across biol. membranes)
    Nucleotides, biological studies
ΙŢ
    Oligonucleotides
    Polynucleotides
     Polysaccharides, biological studies
    Proteins, general, biological studies
     RL: BPR (Biological process); BSU (Biological study, unclassified); PEP
     (Physical, engineering or chemical process); THU (Therapeutic use); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (transport of; conjugates for transporting mols. across biol.
       membranes)
                                               325760-04-1P 325760-05-2P
    89962-57-2P 325760-02-9P 325760-03-0P
IT
    325760-06-3P 325760-07-4P 325760-08-5P 325760-09-6DP, conjugate with
    Cy3 325760-10-9P
    RL: BPR (Biological process); BSU (Biological study, unclassified); PEP
     (Physical, engineering or chemical process); PNU (Preparation,
    unclassified); THU (Therapeutic use); BIOL (Biological study); PREP
     (Preparation); PROC (Process); USES (Uses)
        (conjugates for transporting mols. across biol. membranes)
    146397-20-8D, Cy3, conjugate with oligonucleotides
IT
    RL: BPR (Biological process); BSU (Biological study, unclassified); PEP
     (Physical, engineering or chemical process); THU (Therapeutic use); BIOL
     (Biological study); PROC (Process); USES (Uses)
        (transport of; conjugates for transporting mols. across biol.
       membranes)
    110616-00-7 116364-61-5 146216-12-8 147178-75-4
ΙT
    159845-57-5 161415-79-8 161415-81-2 163665-40-5 164910-61-6
     165447-62-1 166436-80-2 173432-53-6 173432-56-9 173432-57-0
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173432-58-1
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173432-63-8 173432-67-2 173432-68-3 173432-69-4 173432-70-7
173432-71-8 181988-02-3 181988-09-0, 1: PN: WO0004034 SEQID: 1
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AX283174 325605-38-7 325605-39-8 325605-40-1 325605-41-2
325605-42-3 325605-43-4 325605-44-5 325605-45-6 325605-46-7
325605-47-8 325605-48-9 325605-49-0 325605-50-3 325605-51-4
325605-52-5 325761-26-0
RL: PRP (Properties)
   (unclaimed nucleotide sequence; conjugates and methods for the prodn.
  thereof for transporting mols. across biol. membranes)
116364-61-5
RL: PRP (Properties)
   (unclaimed nucleotide sequence; conjugates and methods for the prodn.
  thereof for transporting mols. across biol. membranes)
116364-61-5 HCAPLUS
DNA, d(T-A-T-T-C-C-G-T-C-A-T) (9CI) (CA INDEX NAME)
```

Absolute stereochemistry.

IT

RN

CN

PAGE 1-A

PAGE 2-B

DN 133:89771

TI Olefinic peptide nucleic acids (OPAs): new aspects of the molecular recognition of DNA by PNA

AU Schutz, Rolf; Cantin, Michel; Roberts, Christopher; Greiner, Beate; Uhlmann, Eugen; Leumann, Christian

CS Department of Chemistry and Biochemistry, University of Bern, Bern, 3012, Switz.

SO Angewandte Chemie, International Edition (2000), 39(7), 1250-1253 CODEN: ACIEF5; ISSN: 1433-7851

PB Wiley-VCH Verlag GmbH

DT Journal

LA English

CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 33

GΙ

٦

In order to study the structural and electrostatic effect of the PNA rotameric forms, the authors have synthesized olefinic polyamide nucleic acids (OPAs) in which the central amide functionality was replaced by an isostructural, configurationally stable C-C double bond in either the Z or E configuration (I; BASE = thymidine or adenine), and used them to prep. (E) - or (Z) - OPA oligomers. A series of mono-substituted PNAs and fully-modified (E) and (Z)-OPAs were synthesized and their duplex-forming behavior with DNA studied. Both (E)and (Z)-OPAs bound to complementary DNA with similar affinities as DNA itself, but in contrast to PNA, OPA2/DNA triplexes were not formed, and OPA preferentially bound in the parallel mode to DNA. Results led to the conclusion that amide functionality in the base-linked unit in PNA detd. significantly the affinity and preferred strand orientation in PNA/DNA duplexes, and seemed to be responsible for the propensity to form PNA2/DNA triplexes; these properties do not depend on the conformational constraints that the amide functionality exerts on the base-linker unit, but rather on its electrostatic properties.

ST olefinic peptide nucleic acid PNA

analog prepn hybridization DNA; mol recognition DNA OPA conformation Quaternary structure

(DNA triplex; prepn. and characteristics of olefinic peptide nucleic acids as PNA analogs for mol.

recognition of DNA)

IT Conformation

Nucleic acid hybridization

(prepn. and characteristics of olefinic **peptide nucleic acids** as **PNA** analogs for mol. recognition of DNA)

IT DNA

IT

Nucleic acids

RL: PRP (Properties)

(prepn. and characteristics of olefinic peptide

```
nucleic acids as PNA analogs for mol.
       recognition of DNA)
    Alkenes, preparation
IT
      Peptide nucleic acids
    RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and characteristics of olefinic peptide
       nucleic acids as PNA analogs for mol.
       recognition of DNA)
    161353-44-2P 178036-67-4P 279264-60-7P 279264-61-8P 279264-62-9P
IT
    279694-96-1P 279694-97-2P 280587-99-7P
    RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and characteristics of olefinic peptide
       nucleic acids as PNA analogs for mol.
       recognition of DNA)
    166877-37-8P 226949-23-1P 277322-59-5P 277322-62-0P
IT
    277322-64-2P 277322-66-4P 277322-72-2P
    277322-74-4P 277322-76-6P 277322-77-7P 277322-79-9P
    277322-80-2P 277322-82-4P 279694-94-9P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (prepn. and characteristics of olefinic peptide
       nucleic acids as PNA analogs for mol.
       recognition of DNA)
RE.CNT
             THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD
       22
RE
(1) Almarsson, O; Proc Natl Acad Sci USA 1993, V90, P7518 HCAPLUS
(2) Almarsson, O; Proc Natl Acad Sci USA 1993, V90, P9542 HCAPLUS
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(17) Uhlmann, E; Angew Chem 1996, V108, P2793
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(22) Will, D; Tetrahedron 1995, V51, P12069 HCAPLUS
    277322-64-2P 277322-66-4P 277322-72-2P
IT
    277322-79-9P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (prepn. and characteristics of olefinic peptide
       nucleic acids as PNA analogs for mol.
       recognition of DNA)
    277322-64-2 HCAPLUS
RN
    Peptide nucleic acid, (dT-(5'-deamino-5'-oxy)G[imino[(3Z)-3-[2-(3,4-
CN
    dihydro-5-methyl-2,4-dioxo-1(2H)-pyrimidinyl)ethylidene]-5-oxo-1,5-
    pentanediyl]]A-G-A-T-C-A-C-T)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME)
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Absolute stereochemistry. Double bond geometry as shown.

PAGE 1-A

NH₂

PAGE 1-C

PAGE 2-A

PAGE 2-B

RN 277322-66-4 HCAPLUS

CN Peptide nucleic acid, (dT-(5'-deamino-5'-oxy)G-T-A-G-A[imino[(3Z)-3-[2-(3,4-dihydro-5-methyl-2,4-dioxo-1(2H)-pyrimidinyl)ethylidene]-5-oxo-1,5-pentanediyl]]C-A-C-T)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

PAGE 1-A

PAGE 1-B

PAGE 1-C

PAGE 1-D

PAGE 2-D

||

RN 277322-72-2 HCAPLUS

Peptide nucleic acid, (dT-(5'-deamino-5'-oxy)G-T-A-G-A-T-C-A-C)-[(3E)-5-(3,4-dihydro-5-methyl-2,4-dioxo-1(2H)-pyrimidinyl)-3-[2-[(6-hydroxyhexyl)amino]-2-oxoethyl]-3-pentenyl]NH (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

PAGE 1-A

PAGE 1-B

PAGE 1-D

$$NH_2$$
 NH_2
 NH_2

PAGE 2-B

/ H2N

RN 277322-79-9 HCAPLUS CN Peptide nucleic acid, (dC-T-T-T-T-A-A-T-A)-gly-NH2 (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

PAGE 1-C

PAGE 1-D

L84 ANSWER 5 OF 12 HCAPLUS COPYRIGHT 2003 ACS

AN 2000:127528 HCAPLUS

DN 132:175816

TI Antisense oligonucleotide-based compositions and methods for reducing radiation and drug resistance in cells

IN Chang, Esther H.; Pirollo, Kathleen F.

PA USA

SO U.S., 18 pp. CODEN: USXXAM

DT Patent

LA English

IC ICM C12Q001-68

ICS C12N009-00; C12N015-85; C07H021-04

NCL 435006000

CC 1-6 (Pharmacology)

siew - 09 / 835370 Section cross-reference(s): 8, 63 FAN.CNT 1 APPLICATION NO. DATE PATENT NO. KIND DATE PI US 6027892 A 20000222 US 1997-991830 19971216 PRAI US 1996-34160P P 19961230 Provided are antisense oligonucleotides directed against the raf-1 gene, AB Ha-ras gene and HER-2 gene, components of a signal transduction pathway involving oncogenes and their normal counterparts and leading to the phenotype of cellular radioresistance. Administration of these antisense oligonucleotides is shown to reverse the radioresistance phenotype in cells overexpressing HER-2 or a mutant form of Ha-ras. Methods and compns. for reversing radiation resistance among other conditions involving these genes are disclosed. antisense oligonucleotide drug radiation resistance redn ŞΤ DNA sequences ITDrug resistance Radiation Radiotherapy Signal transduction, biological (antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Antisense oligonucleotides ΙΤ Phosphorothioate oligonucleotides RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) neu (receptor) IT RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Antitumor agents IT(bladder carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Gene, animal IT RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (c-Ha-ras; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Gene, animal ITRL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (c-erbB2; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Gene, animal ITRL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (c-raf-1; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Bladder ITBladder Head Head Lung, neoplasm

Mammary gland
Mammary gland
Neck, anatomical
Neck, anatomical
Ovary, neoplasm

Lung, neoplasm

Ovary, neoplasm Pancreas, neoplasm Pancreas, neoplasm Prostate gland Prostate gland Stomach, neoplasm Stomach, neoplasm (carcinoma, inhibitors; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Bladder Head Lung, neoplasm Mammary gland Neck, anatomical Ovary, neoplasm Pancreas, neoplasm Prostate gland Stomach, neoplasm (carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Antitumor agents (cervix carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Uterus, neoplasm Uterus, neoplasm (cervix, carcinoma, inhibitors; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells) Uterus, neoplasm (cervix, carcinoma; antisense oligonucleotide-based compns. and methods Antitumor agents Intestine, neoplasm Intestine, neoplasm

for reducing radiation and drug resistance in cells)

(colon carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

IT

IT

 IT

IT

IT

(colon, carcinoma, inhibitors; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Intestine, neoplasm IT

> (colon, carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Antitumor agents IT

(head and neck squamous cell carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Antitumor agents IT

> (head carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Drug delivery systems IT

(liposomes; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Antitumor agents IT

> (lung carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Antitumor agents IT

> (mammary gland carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Antitumor agents IT

> (neck carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Antitumor agents IT

(ovary carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

Antitumor agents IT

(pancreas carcinoma; antisense oligonucleotide-based compns. and
 methods for reducing radiation and drug resistance in cells)
Antitumor agents
 (prostate carcinoma; antisense oligonucleotide-based compns. and
 methods for reducing radiation and drug resistance in cells)
Head
Head

IT Head Head

ΙT

Neck, anatomical

Neck, anatomical

(squamous cell carcinoma, inhibitors; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

IT Head

Neck, anatomical

(squamous cell carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

IT Antitumor agents

(stomach carcinoma; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

IT **116364-61-5** 125486-19-3 158768-80-0 259113-38-7 259158-20-8 259158-21-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

IT 139691-76-2

RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

IT 259168-28-0, 4: PN: US6027892 SEQID: 4 unclaimed DNA 259168-29-1, 5: PN: US6027892 SEQID: 5 unclaimed DNA 259168-30-4, 6: PN: US6027892 SEQID: 6 unclaimed DNA 259168-31-5, 8: PN: US6027892 SEQID: 8 unclaimed DNA 259168-32-6, 9: PN: US6027892 SEQID: 9 unclaimed DNA RL: PRP (Properties)

(unclaimed nucleotide sequence; antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

RE.CNT 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD RE

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- (25) Tseng; Cancer Gene Therapy 1994, V1(1), P65 HCAPLUS
- (26) Vaughn; Nucleic Acids Res 1996, V24, P4558 HCAPLUS
- IT 116364-61-5

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(antisense oligonucleotide-based compns. and methods for reducing radiation and drug resistance in cells)

RN 116364-61-5 HCAPLUS

CN DNA, d(T-A-T-T-C-C-G-T-C-A-T) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 2-B

PAGE 3-B

```
Correction of: 1996:755988
    127:2136
DN
      Correction of: 126:141081
     Synthesis and properties of PNA/DNA chimeras
TI
ΑU
    Uhlmann, Eugen; Will, David W.; Breipohl,
     Gerhard; Langner, Dietrich; Ryte, Antonina
     Hoechst AG, Frankfurt/Main, D-65926, Germany
CS
SO
     Angewandte Chemie, International Edition in English (1996), 35(22),
     2632-2635
    CODEN: ACIEAY; ISSN: 0570-0833
    VCH
PB
     Journal
\mathsf{DT}
    English
LA
CC
    6-2 (General Biochemistry)
     Section cross-reference(s): 3, 9
    We have developed a generally applicable method for the automated
AB
     synthesis of DNA/PNA chimeras. This method is fully compatible
     with std. DNA synthesis methods and requires no addnl. deprotection steps
     at the end of oligomer synthesis. The binding affinity of DNA-PNA
     chimeras is higher than that of the comparable DNA-phosphorothioate
     chimeras or natural oligonucleotides. Unlike pure PNAs, the
     DNA-PNA chimeras investigated bind only in the antiparallel
     orientation to their complementary nucleic acids under physiol conditions.
    PNA DNA chimera prepn automated
ST
    104655-85-8 149376-29-4 170490-73-0 172316-36-8 172316-40-4
ΙT
    172316-41-5 172316-42-6 185810-72-4 185810-73-5 185810-74-6
                 185810-78-0 185810-79-1 185810-80-4 185810-81-5
    185810-76-8
    185810-82-6
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reactant in synthesis of PNA/DNA chimeras)
    172316-39-1 185831-40-7 185831-41-8 185831-42-9
    185831-43-0 185831-44-1 185970-57-4 185970-58-5 185970-59-6
    185970-60-9 185970-61-0 185970-62-1 186050-47-5 186050-48-6
    186050-49-7 186050-51-1 186050-52-2 186050-53-3
    186050-54-4 186050-55-5 186050-56-6 186050-57-7 186050-58-8
     RL: BPR (Biological process); BSU (Biological study, unclassified); PEP
     (Physical, engineering or chemical process); PRP (Properties); BIOL
     (Biological study); PROC (Process)
        (synthesis and properties of PNA/DNA chimeras)
    186050-42-0P 186050-43-1P 186050-44-2P 186050-45-3P
                                                               186050-46-4P
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (synthesis and properties of PNA/DNA chimeras)
    172316-39-1 186050-51-1
ΙT
     RL: BPR (Biological process); BSU (Biological study, unclassified); PEP
     (Physical, engineering or chemical process); PRP (Properties); BIOL
     (Biological study); PROC (Process)
        (synthesis and properties of PNA/DNA chimeras)
     172316-39-1 HCAPLUS
RN
     Peptide nucleic acid, (H-A-C-A-T-C-A-T-G-G-T-C-G)-(6-hydroxyhexyl)NH (9CI)
CN
       (CA INDEX NAME)
```

PAGE 1-A

PAGE 2-A

PAGE 3-B

PAGE 3-C

PAGE 4-D

RN 186050-51-1 HCAPLUS

CN DNA, d(T-A-T-T-C-C-G-T-C-A-T), complex with peptide nucleic acid (dA-dT-dG-(5'-deamino-5'-oxy)A-C-G-G-A-A-T-A)-(6-hydroxyhexyl)NH (1:1) (9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

L84 ANSWER 7 OF 12 HCAPLUS COPYRIGHT 2003 ACS

AN 1996:755988 HCAPLUS

DN 126:141081

TI Synthesis and properties of PNA/DNA chimeras

AU Uhlmann, Eugen; Will, David W.; Breiphohl, Gerhard; Langner, Dietrich; Ryte, Antonina

CS Hoechst AG, Frankfurt/Main, D-65926, Germany

SO Angewandte Chemie, International Edition in English (1996), 35(22), 2632-2635

CODEN: ACIEAY; ISSN: 0570-0833

PB VCH

DT Journal

LA English

CC 6-2 (General Biochemistry)
Section cross-reference(s): 32, 33

AB We have developed a generally applicable method for the automated synthesis of DNA/PNA chimeras. This method is fully compatible with std. DNA synthesis methods and requires no addnl. deprotection steps at the end of oligomer synthesis. The binding affinity of DNA-PNA chimeras is higher than that of the comparable DNA-phosphorothicate chimeras or natural oligonucleotides. Unlike pure PNAs, the

Chimeras or natural oligonucleotides. Unlike pure PNAS, the DNA-PNA chimeras investigated bind only in the antiparallel orientation to their complementary nucleic acids under physiol.

conditions.

ST PNA DNA chimera prepn automated

104655-85-8 149376-29-4 170490-73-0 172316-36-8 ΙT 172316-40-4 172316-41-5 172316-42-6 185810-72-4 185810-73-5 185810-74-6 185810-78-0 185810-79-1 185810-80-4 185810-81-5 185810-76-8 185810-82-6 RL: RCT (Reactant); RACT (Reactant or reagent) (reactant in synthesis of PNA/DNA chimeras) **172316-39-1** 185831-40-7 185831-41-8 185831-42-9 IT 185831-43-0 185831-44-1 185970-57-4 185970-58-5 185970-59**-**6 185970-60-9 185970-61-0 185970-62-1 186050-47-5 186050-48-6 186050-49-7 **186050-51-1** 186050-52-2 186050-53-3 186050-54-4 186050-55-5 186050-56-6 186050-57-7 186050-58-8 RL: BPR (Biological process); BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); PRP (Properties); BIOL (Biological study); PROC (Process) (synthesis and properties of PNA/DNA chimeras) 186050-42-0P 186050-43-1P 186050-44-2P 186050-45-3P IT186050-46-4P RL: SPN (Synthetic preparation); PREP (Preparation) (synthesis and properties of PNA/DNA chimeras) 172316-39-1 186050-51-1 ITRL: BPR (Biological process); BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); PRP (Properties); BIOL (Biological study); PROC (Process) (synthesis and properties of PNA/DNA chimeras) 172316-39-1 HCAPLUS RNPeptide nucleic acid, (H-A-C-A-T-C-A-T-G-G-T-C-G)-(6-hydroxyhexyl)NH (9CI) CN(CA INDEX NAME)

PAGE 1-A

PAGE 2-A

PAGE 3-A

PAGE 3-B

PAGE 3-C

PAGE 4-D

RN 186050-51-1 HCAPLUS
CN DNA, d(T-A-T-T-C-C-G-T-C-A-T), complex with peptide nucleic acid
(dA-dT-dG-(5'-deamino-5'-oxy)A-C-G-G-A-A-T-A)-(6-hydroxyhexyl)NH (1:1)
(9CI) (CA INDEX NAME)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

```
ANSWER 8 OF 12 HCAPLUS COPYRIGHT 2003 ACS
 L84
     1995:994428 HCAPLUS
 ΑN
     124:87805
 DN
     Peptide nucleic acid synthesis using an
 TI
      amino protecting group which is labile to weak acids.
     Breipohl, Gerhard Dr; Uhlmann, Eugen Dr
 IN
     Hoechst A.-G., Germany
 PΑ
      Eur. Pat. Appl., 19 pp.
 SO
     CODEN: EPXXDW
     Patent
 \mathsf{DT}
 LA
     German
      ICM C08G069-06
 IC
      ICS C07D239-54; C07D239-46; C07D473-18; C07D473-34; C08G069-10
      34-3 (Amino Acids, Peptides, and Proteins)
 CC
      Section cross-reference(s): 33
 FAN.CNT 1
      PATENT NO. KIND DATE
                                         APPLICATION NO. DATE
     EP 672700 A1 19950920
                                         EP 1995-103318 19950308
. PI
     EP 672700 B1 19990602
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
      DE 4408531 A1 19950928 DE 1994-4408531 19940314
     AT 180805 E 19990615 AT 1995-103318 19950308
     ES 2132450 T3 19990816
                                        ES 1995-103318 19950308
                                        FI 1995-1130 19950310
     FI 9501130 A 19950915
     AU 9514801 A1 19950921
                                         AU 1995-14801
                                                        19950310
     AU 695931 B2 19980827
     CA 2144477 AA 19950915
                                         CA 1995-2144477 19950313
                                         NO 1995-957 19950313
     NO 9500957 A 19950915
                                         JP 1995-54642
      JP 07285989 A2 19951031
                                                        19950314
     US 6046306
                           20000404
                                         US 1997-927178
                                                         19970911
 PRAI DE 1994-4408531
                          19940314
                          19950313
     US 1995-402385
      RAk(XB1)nQlQ1 [XB = NH(CH2)fCH2N(COCH2B)(CH2)fO, NHCH[(CH2)fB]CONHCH2CO,
 AΒ
      NHCH[(CH2)fB](CH2)3CO, etc.; f = 1-4; k, l = 0-10; A, Q = amino acid
      residue; B = (un)natural nucleic acid base or prodrug or replacement forms
      thereof; Q1 = OH, amino], were prepd. by solid phase synthesis. Thus,
      H-[Aeg(T)]3hex [Aeg(T) = N-(2-aminoethyl)-N-[(1-thyminyl)acetyl]glycyl,
      hex = HN(CH2)6OH] was prepd. on hex-succ-tentagel (succ = succinoyl)
      (prepn. given) on a DNA synthesizer.
      peptide nucleic acid synthesis protecting
 ST
      group
      Protective groups
 IT
         (peptide nucleic acid synthesis using an
        amino protecting group which is labile to weak acids)
     Nucleopeptides
 IT
      RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
      (Preparation)
         (peptide nucleic acid synthesis using an
        amino protecting group which is labile to weak acids)
      172316-43-7P
 IT
      RL: BYP (Byproduct); PREP (Preparation)
         (peptide nucleic acid synthesis using an
        amino protecting group which is labile to weak acids)
      172316-37-9P 172316-38-0P 172316-39-1P
 ΙT
      RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
      (Preparation)
         (peptide nucleic acid synthesis using an
        amino protecting group which is labile to weak acids)
      108-30-5, reactions 4048-33-3, 6-Amino-1-hexanol 14470-28-1
 IT
      172316-36-8 172316-40-4 172316-41-5 172316-42-6 172316-44-8
      172316-45-9
      RL: RCT (Reactant); RACT (Reactant or reagent)
```

(peptide nucleic acid synthesis using an

amino protecting group which is labile to weak acids)

IT 114729-83-8P 172316-34-6P 172316-35-7DP, resin-bound

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(peptide nucleic acid synthesis using an

amino protecting group which is labile to weak acids)

IT 172316-39-1P

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(peptide nucleic acid synthesis using an

amino protecting group which is labile to weak acids)

RN 172316-39-1 HCAPLUS

CN Peptide nucleic acid, (H-A-C-A-T-C-A-T-G-G-T-C-G)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

PAGE 3-A

PAGE 3-B

PAGE 3-C

PAGE 4-D

$$\begin{array}{c|c} CH_2-CH_2-NH_2 \\ -N-C=0 \\ CH_2 \\ N \\ N \\ NH_2 \end{array}$$

H2N

L84 ANSWER 9 OF 12 HCAPLUS COPYRIGHT 2003 ACS

AN 1995:994427 HCAPLUS

DN 124:87804

TI Peptide nucleic acid synthesis using a base labile amino protecting group.

IN Breipohl, Gerhard Dr; Uhlmann, Eugen Dr; Knolle,

```
Jochen Dr
    Hoechst A.-G., Germany
PA
    Eur. Pat. Appl., 31 pp.
SO
    CODEN: EPXXDW
    Patent
DT
LA
    German
    ICM C08G069-06
IC
    ICS C07D473-18; C07D473-34; C07D239-54; C07D239-46; C08G069-10
    34-3 (Amino Acids, Peptides, and Proteins)
CC
    Section cross-reference(s): 33
FAN.CNT 1
                                       APPLICATION NO.
                                                       DATE
    PATENT NO. KIND DATE
    EP 672701 A1 19950920
                                                      19950308
                                       EP 1995-103319
PI
    EP 672701 B1 19990728
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
    DE 4408533 A1 19950928 DE 1994-4408533 19940314
    AT 182602 E 19990815 AT 1995-103319 19950308
    ES 2136755 T3 19991201
                                       ES 1995-103319 19950308
    FI 9501129 A 19950915
                                       FI 1995-1129 19950310
                                       AU 1995-14800
                                                      19950310
    AU 9514800 A1 19950921
    AU 683714 B2
                       19971120
    CA 2144473
                    AA 19950915
                                       CA 1995-2144473 19950313
                                       NO 1995-958 19950313
    NO 9500958
                         19950915
                     A
                                       JP 1995-54641 19950314
    JP 07291909
                     Α2
                        19951107
                                       US 1997-967197 19971029
    US 6121418
                     A
                          20000919
                                       US 2000-495457
    US 6316595
                                                       20000201
                        20011113
               B1
PRAI DE 1994-4408533 A 19940314
                     В1
    US 1995-402844
                        19950313
    US 1997-967197
                     Α3
                         19971029
    RAk[NHCH2CH2N(COCH2B)CH2CO]nQlQ1 (R = H, alkanoyl, alkoxycarbonyl,
    cycloalkanoyl, aroyl, heteroaroyl, group which promotes intracellular
    uptake or interacts with target nucleic acids; A, Q = amino acid residue;
    Q1 = OH, amino; B = nucleobase or prodrug form thereof; l = 0-20; n = 0
    1-50), were prepd. by solid phase synthesis. Thus, H-[Aeg(T)]8-Lys-NH2 [
    Aeg(T) = N-(2-aminoethyl)-N-[(1-thyminyl)acetyl]glycyl] was prepd. by
    coupling of FMOC-Lys(BOC)-OH and FMOC-Aeg(T)-OH (prepn. given) on
    5-(FMOC-amino-4-methoxybenzyl)-2,4-dimethoxyphenylpropionic
    acid-derivatized aminomethylpolystyrene resin using an activator soln. of
    PyBOP (PyBOP = benzotriazolyl-1-oxytripyrrolidiniophosphonium
    hexafluorophosphate) in DMF, NEM (N-ethylmorpholine) in DMF as base for
    activation, and 20% piperidine in DMF for deprotection.
    peptide nucleic acid synthesis base labile;
ST
    base labile protecting group pna synthesis
    Merrifield synthesis
IT
       (peptide nucleic acid synthesis using a
       base labile amino protecting group)
    Nucleopeptides
IT
    RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
    (Preparation)
       (peptide nucleic acid synthesis using a
       base labile amino protecting group)
    139166-84-0P 172405-66-2P 172405-67-3P 172405-68-4P
IT
    172405-69-5P
    RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
    (Preparation)
       (peptide nucleic acid synthesis using a
       base labile amino protecting group)
    65-71-4, Thymine 71-30-7, Cytosine 73-24-5, 6-Aminopurine, reactions
\operatorname{IT}
    96-32-2, Methyl bromoacetate 108-53-2 10310-21-1, 2-Amino-6-
    chloropurine 18907-79-4 20924-05-4 24123-14-6 71989-14-5
    71989-26-9
    RL: RCT (Reactant); RACT (Reactant or reagent)
```

(peptide nucleic acid synthesis using a

base labile amino protecting group)

 IT
 13251-16-6P
 55036-34-5P
 67826-12-4P
 119451-90-0P
 169396-92-3P

 172405-14-0P
 172405-15-1P
 172405-16-2P
 172405-27-5P
 172405-43-5P

 172405-44-6P
 172405-45-7P
 172405-46-8P
 172405-47-9P
 172405-48-0P

 172405-54-8P
 172405-55-9P
 172405-56-0P
 172405-57-1P
 172405-58-2P

 172405-64-0P
 172405-65-1P
 172405-61-7P
 172405-62-8P
 172405-63-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(peptide nucleic acid synthesis using a

base labile amino protecting group)

IT 172405-68-4P 172405-69-5P

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(peptide nucleic acid synthesis using a

base labile amino protecting group)

RN 172405-68-4 HCAPLUS

CN Peptide nucleic acid, (acetyl-A-C-A-T-C-A-T-G-G-T-C-G)-Lys-NH2 (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

PAGE 1-C

PAGE 1-D

PAGE 1-E

__NHAc

RN 172405-69-5 HCAPLUS
CN Peptide nucleic acid, (Asp-C-C-A-T-G-G-T-C-C-C)-Asp-[N-(6-hydroxyhexyl)]NH (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

PAGE 1-C

PAGE 1-D

PAGE 2-A

PAGE 2-C

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L84 ANSWER 10 OF 12 HCAPLUS COPYRIGHT 2003 ACS
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AN 1995:908968 HCAPLUS

DN 124:117857

TI The synthesis of polyamide nucleic acids using a novel monomethoxytrityl protecting-group strategy

AU Will, David W.; Breipohl, Gerhard; Langner, Dietrich; Knolle, Jochen; Uhlmann, Eugen

CS Hoechst AG, Allgemeine Pharma Forschung G838, Frankfurt am Main, D-65926, Germany

SO Tetrahedron (1995), 51(44), 12069-82 CODEN: TETRAB; ISSN: 0040-4020

PB Elsevier

DT Journal

LA English

CC 33-9 (Carbohydrates)

OS CASREACT 124:117857

The prepn. of 4-MeOC6H4CPh2NHCH2CH2N(COCH2R)CH2CO2Me (R = thymine, N4-tert-butylbenzoylcytosine, N6-anisoyladenine, N2-isobutanoylguanine) for the synthesis of polyamide nucleic acids (PNAs) is described. The use of base-labile acyl-type nucleobase protecting groups, including monomethyltrityl N-protection of H2NCH2CH2NhCH2CO2Me, and of a succinyl-linked solid-support offers a synthetic strategy similar to std. oligonucleotide synthesis conditions. This strategy has been successfully applied for the synthesis of PNAs of mixed base sequence.

ST polyamide nucleic acid analog prepn; monomethoxytrityl amine protecting group aminoethylglycine; solid phase synthesis polyamide oligonucleotide analog

IT Nucleic acids

RL: SPN (Synthetic preparation); PREP (Preparation) (analogs, synthesis of polyamide nucleic acid analogs from monomethoxytrityl-protected aminoethylglycine)

IT Protective groups

(methoxytrityl, for amine in aminoethylglycine)

IT 71-30-7, Cytosine 73-24-5, Adenine, reactions 73-40-5 96-32-2,
 Methyl bromoacetate 107-15-3, 1,2-Ethanediamine, reactions 298-12-4
 1710-98-1, 4-tert-Butylbenzoyl chloride 4048-33-3, 6-Aminohexan-1-ol
 20924-05-4

RL: RCT (Reactant); RACT (Reactant or reagent)

(synthesis of polyamide nucleic acid analogs from monomethoxytrityl-protected aminoethylglycine)

1T 18907-79-4P 21047-89-2P 24123-14-6P, N-(2-Aminoethyl)glycine 97025-97-3P 114729-83-8P 135697-25-5P 170944-06-6P 172316-34-6DP, polymer bound 172316-34-6DP, polymer-bound 172316-34-6P 172316-36-8P 172316-40-4P 172316-42-6P 172316-45-9P 172405-11-7P 172405-12-8P 172405-17-3P 172405-18-4P 172405-19-5P 172405-20-8P 172405-21-9P 172405-39-9P 172405-41-3P 172405-42-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis of polyamide nucleic acid analogs from monomethoxytrityl-protected aminoethylglycine)

IT 172316-39-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (synthesis of polyamide nucleic acid analogs from monomethoxytrityl-protected aminoethylglycine)

IT 172316-39-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (synthesis of polyamide nucleic acid analogs from monomethoxytrityl-protected aminoethylglycine)

RN 172316-39-1 HCAPLUS

CN Peptide nucleic acid, (H-A-C-A-T-C-A-T-G-G-T-C-G)-(6-hydroxyhexyl)NH (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 3-A

PAGE 3-B

$$NH_2$$
 NH_2
 NH_2

PAGE 3-C

PAGE 4-D

L84 ANSWER 11 OF 12 HCAPLUS COPYRIGHT 2003 ACS

AN 1990:111614 HCAPLUS

DN 112:111614

TI Comparative inhibition of ras p21 protein synthesis with phosphorus-modified antisense oligonucleotides

AU Chang, E. H.; Yu, Z.; Shinozuka, K.; Zon, G.; Wilson, W. D.; Strekowska,

```
Α.
     Dep. Pathol., Uniformed Serv. Univ. Health Sci., Bethesda, MD, 20814, USA
CS
     Anti-Cancer Drug Design (1989), 4(3), 221-32
SO
     CODEN: ACDDEA; ISSN: 0266-9536
     Journal
DT
     English
LA
CC
    1-6 (Pharmacology)
    A rabbit reticulocyte lysate translation assay was used to quant. compare
AB
     a series of antisense oligodeoxyribonucleotides (11-mers) having different
     internucleoside linkages and various degrees of complementarity (100-80%)
     with the start codon and downstream 8 bases of Balb-ras p21 mRNA.
     oligomers had contiguous phosphodiester, alternating methylphosphonate-
     phosphodiester, contiguous methylphosphonate, or contiguous
     phosphorothioate linkages. The test compds. present in
     .apprx.103-104-fold excess over mRNA (15 nM mRNA) inhibited protein
     synthesis to a degree which was dependent on the concn. and the oligomer
     sequence. At low concns. (12.5-25 .mu.M), the phosphorothioate analogs
     were the most potent inhibitors of p21 protein synthesis but the
     sequence-nonspecific effect for these oligomers was dominant at higher
     concns. (100-200 .mu.M). The methylphosphonate oligomers were slightly
     more discriminant. Relative hybridization strengths were assessed by
     melting studies using a DNA oligomer target to mimic the mRNA.
     oligodeoxyribonucleotide antisense ras p21 protein synthesis; antitumor
ST
     oligodeoxyribonucleotide antisense p21 protein synthesis
     Neoplasm inhibitors
IT
        (antisense oliqodeoxyribonucleotides inhibition of ras p21 protein
        synthesis in relation to)
     Protein formation
IT
        (of ras p21, antisense oligodeoxyribonucleotides inhibition of)
     Nucleotides, polymers
IT
     RL: BIOL (Biological study)
        (oligo-, deoxyribo-, protein ras p21 formation inhibition by antisense)
     116338-84-2 116364-61-5 124306-02-1 124306-03-2
IT
    124306-04-3 125486-17-1 125486-18-2 125486-19-3
                                                            125486-20-6
     125486-21-7 125486-22-8 125486-23-9 125500-19-8
     RL: BIOL (Biological study)
        (protein ras p21 formation inhibition by)
     116364-61-5
IT
     RL: BIOL (Biological study)
        (protein ras p21 formation inhibition by)
     116364-61-5 HCAPLUS
RN
     DNA, d(T-A-T-T-C-C-G-T-C-A-T) (9CI) (CA INDEX NAME)
CN
```

Absolute stereochemistry.

PAGE 1-A

PAGE 2-B

PAGE 3-B

L84 ANSWER 12 OF 12 HCAPLUS COPYRIGHT 2003 ACS

AN 1988:524551 HCAPLUS

DN 109:124551

TI Unusual duplex formation in purine rich oligodeoxyribonucleotides

AU Wilson, W. David; Do Trong Minh Hoa; Zuo, Elizabeth T.; Zon, Gerald

CS Dep. Chem., Georgia State Univ., Atlanta, GA, 30303-3083, USA

SO Nucleic Acids Research (1988), 16(11), 5137-51

CODEN: NARHAD; ISSN: 0305-1048

DT Journal

LA English

CC 6-2 (General Biochemistry)

The purine rich oligodeoxyribonucleotides 1C, [d(ATGACGGAATA)], and 2C, AΒ [d(ATGAGCGAATA)], alone exhibit highly cooperative melting transitions. Anal. of the concn. dependence of melting, and electrophoretic studies indicate that these oligomers can form an unusual purine rich offset double helix. The unusual duplex is predicted to contain 4 A.cntdot.T, 2 G.cntdot.C, and 4 G.cntdot.A mismatch base pairs as well as a single A base stacked on the 3' end of each chain of the helix. Other possible models for the duplex are unlikely because they are predicted to contain many base pairs of low stability. Changing the central sequence to CGG or GGG should destabilize the duplex and this is obsd. The unusual duplex of 2C is more stable than the duplex of 1C, indicating that the stability of G.cntdot.A base pairs is quite sensitive to the surrounding sequence. Addn. of 1C and 2C to their complementary pyrimidine strands results in normal duplexes of similar stability. Apparently, the unusual duplexes are significantly stabilized by the intrinsic stacking tendency of purine bases.

ST oligodeoxyribonucleotide duplex formation purine rich; DNA oligodeoxyribonucleotide duplex formation

IT Entropy Free energy Thermodynamics

(of duplex formation by oligodeoxyribonucleotides, unusual purine offset double helix formation in relation to)

IT Heat of formation

(of duplex formation in oligodeoxyribonucleotides, unusual purine offset double helix formation in relation to)

IT Quaternary structure

(of purine rich oligodeoxyribonucleotides)

IT Nucleotides, polymers

RL: BIOL (Biological study)

(oligo-, deoxyribo-, offset double helix formation by purine rich)

IT 73-40-5, Guanine

RL: BIOL (Biological study)

(adenine base pair with, in unusual duplex in purine rich oligodeoxynucleotide)

IT 73-24-5, Adenine, biological studies

RL: BIOL (Biological study)

(guanine base pair with, in unusual duplex of purine rich oligodeoxynucleotides)

116338-84-2 **116364-61-5** 116364-62-6 116364-63-7

116364-64-8 116374-13-1

RL: BIOL (Biological study)

(melting curve of, unusual purine rich offset double helix formation in relation to)

IT 116338-85-3 116338-86-4

RL: BIOL (Biological study)

(self-complementary duplex formation by, unusual purine rich offset double helix formation in)

IT 116364-61-5

IT

RL: BIOL (Biological study)

(melting curve of, unusual purine rich offset double helix formation in relation to)

RN 116364-61-5 HCAPLUS

CN DNA, d(T-A-T-T-C-C-G-T-C-A-T) (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 2-B

PAGE 3-B

L85 26 L72 NOT (L69 OR L84)

=> d all tot 185

L85 ANSWER 1 OF 26 HCAPLUS COPYRIGHT 2003 ACS

AN 2002:805617 HCAPLUS

- TI (2'-O-methyl-RNA)-3'-PNA chimeras: A new class of mixed backbone oligonucleotide analogues with high binding affinity to RNA
- AU Greiner, Beate; Breipohl, Gerhard; Uhlmann, Eugen
- CS Aventis Pharma Deutschland GmbH, Frankfurt a.M., D-65926, Germany
- SO Helvetica Chimica Acta (2002), 85(9), 2619-2626 CODEN: HCACAV; ISSN: 0018-019X
- PB Verlag Helvetica Chimica Acta
- DT Journal
- LA English
- CC 63 (Pharmaceuticals)
- The automated online synthesis of DNA-3'-PNA chimeras 1-4 and AB(2'-O-methyl-RNA)-3'-PNA chimeras 5-8 is described, in which the 3'-terminal part of the oligonucleotide is linked to the N-terminal part of the PNA via N-(.omega.-hydroxyalkyl)-N-[(thymin-1yl)acetyl]qlycine units (alkyl=Et, Ph, Bu, and pentyl). By means of UV thermal denaturation, the binding affinities of all chimeras were directly compared by detg. their Tm values in the duplex with complementary DNA and RNA. All investigated DNA-3'-PNA chimeras and (2'-O-methyl-RNA)-3'-PNA chimeras form more-stable duplexes with complementary DNA and RNA than the corresponding unmodified DNA. Interestingly, a N-(3-hydroxypropyl)glycine linker resulted in the highest binding affinity for DNA-3'-PNA chimeras, whereas the (2'-O-methyl-RNA)-3'-PNA chimeras showed optimal binding with the homologous N-(4-hydroxybutyl)glycine linker. The duplexes of (2'-O-methyl-RNA)-3'-PNA chimeras and RNA were significantly more stable than those contg. the corresponding DNA-3'-PNA chimeras. Surprisingly, we found that the charged (2'-0-methyl-RNA)-3'-PNA chimera with a N-(4-hydroxybutyl)glycine-based unit at the junction to the PNA part shows the same binding affinity to RNA as uncharged PNA. Potential applications of (2'-O-methyl-RNA)-3'-PNA chimeras include their use as antisense agents acting by a RNase-independent mechanism of action, a prerequisite for antisense-oligonucleotide-mediated correction of aberrant splicing of pre-mRNA.
- RE.CNT 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD RE
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- L85 ANSWER 2 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 2001:748116 HCAPLUS
- DN 135:269645

```
Method and device for detecting molecules by means of impedance
\mathrm{T}\,\mathrm{I}
     spectroscopy
     Escher, Claus; Windhab, Norbert; Muth, Jochen
IN
    Aventis Research & Technologies Gmbh & Co. K.-G., Germany
PA
     PCT Int. Appl., 20 pp.
SO
     CODEN: PIXXD2
     Patent
DT
    German
LA
     ICM G01N033-543
IC
     ICS C12Q001-00; C12Q001-68; G01N027-327
     9-7 (Biochemical Methods)
CC
FAN.CNT 1
     PATENT NO. KIND DATE
                                          APPLICATION NO. DATE
                                          WO 2001-EP1899 20010220
    WO 2001075445 A1 20011011
PΙ
        W: JP, US
        RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
            PT, SE, TR
     DE 10015547 A1 20011031
                                          DE 2000-10015547 20000330
     DE 10015547 C2 20020214
     EP 1272851 A1 20030108 EP 2001-929349 20010220
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI, CY, TR
PRAI DE 2000-10015547 A
                           20000330
     WO 2001-EP1899 W
                           20010220
    A method for detecting target structures is characterized in that a
AB
     three-dimensional porous support consisting of a non-conducting material
     is provided with a soln. contg. mols. to be detected and the extent to
     which the support is charged with these mols. is then detd. by measuring
     elec. impedance. The invention also relates to a device for detecting
    target structures by means of impedance measurement. This device consists
     of a chip with a layered structure contg. at least one layer that contains
     electrodes which can be switched in relation to each other, and the porous
     support consists of the non-conducting material, which is placed on
     and(or) under this layer. Thus, oligonucleotides may be used with a nylon
     membrane support.
     impedance spectroscopy biochip electrode
ST
     Proteins, specific or class
{	t IT}
     RL: ARG (Analytical reagent use); DEV (Device component use); ANST
     (Analytical study); USES (Uses)
        (DNA-binding; method and device for detecting mols. by means of
        impedance spectroscopy)
     Biotechnology
{
m IT}
        (biochips; method and device for detecting mols, by means of impedance
        spectroscopy)
    Nucleic acids
\operatorname{IT}
     RL: ARG (Analytical reagent use); DEV (Device component use); ANST
     (Analytical study); USES (Uses)
        (fragments; method and device for detecting mols. by means of impedance
        spectroscopy)
     Enzymes, uses
IT
     RL: ARG (Analytical reagent use); DEV (Device component use); ANST
     (Analytical study); USES (Uses)
        (inhibitors and activators; method and device for detecting mols. by
        means of impedance spectroscopy)
     Fluoropolymers, uses
IT
     Polyamides, uses
     RL: DEV (Device component use); USES (Uses)
        (membrane; method and device for detecting mols. by means of impedance
        spectroscopy)
     Biosensors
IT
```

Electric impedance

Electrodes

(method and device for detecting mols. by means of impedance spectroscopy) Coenzymes ΙT Enzymes, uses Oligonucleotides Peptide nucleic acids Peptides, uses Proteins, general, uses Receptors Transcription factors cDNA RL: ARG (Analytical reagent use); DEV (Device component use); ANST (Analytical study); USES (Uses) (method and device for detecting mols. by means of impedance spectroscopy) Membranes, nonbiological IT(supports; method and device for detecting mols. by means of impedance spectroscopy) 9003-07-0, Polypropylene 9004-35-7, Cellulose acetate ΙT 9004-70-0, Nitrocellulose 24937-79-9, PVDF RL: DEV (Device component use); USES (Uses) (membrane; method and device for detecting mols. by means of impedance spectroscopy) THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 7 RE (1) Australian Membrane And Biotechnology Research Institute; WO 9744651 A 1997 HCAPLUS (2) Cambridge Life Sciences Plc; WO 9428414 A 1994 HCAPLUS (3) Commissariat A L'Energie Atomique Etabliss de Caract Scient Tech; FR 2757949 A 1998 HCAPLUS (4) Innogenetics N V; WO 9721094 A 1997 HCAPLUS (5) Stetter, J; US 5567301 A 1996 HCAPLUS (6) The John Hopkins University; WO 8809499 A 1988 HCAPLUS (7) The Victoria University Of Manchester; WO 9819153 A 1998 HCAPLUS ANSWER 3 OF 26 HCAPLUS COPYRIGHT 2003 ACS L85 2001:747816 HCAPLUS AN135:302893 DNImmunogenic peptides derived from prostate-specific membrane antigen TI(PSMA) and uses thereof Pedyczak, Arthur; Chong, Pele; Sia, Charles Dwo Yuan ΙN PAAventis Pasteur Limited, Can. SO PCT Int. Appl., 47 pp. CODEN: PIXXD2 Patent DTEnglish LΑ ICICM C07K007-00 ÇÇ 15-2 (Immunochemistry) FAN.CNT 1 PATENT NO. APPLICATION NO. DATE KIND DATE WO 2001074845 A2 20011011 WO 2001-CA411 20010330 PIWO 2001074845 A3 20020510 AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,

DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,

20030206 US 2001-821734 20010330

BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

US 2003027246 A1

PRAI US 2000-193386P P 20000331 The identification of immunogenic peptides of PSMA, nucleic acids coding therefor, and recombinant nucleic acids into which are inserted said nucleic acids coding for PSMA peptides are disclosed. These peptides, nucleic acids and recombinant nucleic acids may be used in isolation, or as compns. thereof to modulate immune responses in animals. The invention further encompasses methods per se of modulating immune responses in animals. vaccine prostate cancer PMSA peptide cytotoxic T lymphocyte STHistocompatibility antigens ITRL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (HLA, class I; immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Histocompatibility antigens IT RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (HLA-A; immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Fusion proteins (chimeric proteins) ΙT RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (PMSA peptide-contg.; immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Immunostimulants IT(adjuvants; immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) T cell (lymphocyte) IT(cytotoxic; immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Oligodeoxyribonucleotides ΙT Oligonucleotides RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (encoding PMSA peptide; immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Antitumor agents ITGenetic vectors Immunity Transformation, genetic (immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Peptides, biological studies ITRL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); PRP (Properties); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Prostate-specific antigen ITRL: BSU (Biological study, unclassified); BIOL (Biological study) (immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Prostate gland ΙΤ (neoplasm, inhibitors; immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer) Antitumor agents IT (prostate gland; immunogenic peptides derived from prostate-specific membrane antigen (PSMA) and uses in treatment of prostate cancer)

Vaccines

 IT

```
(tumor; immunogenic peptides derived from prostate-specific membrane
        antigen (PSMA) and uses in treatment of prostate cancer)
    Antitumor agents
ΙT
        (vaccines; immunogenic peptides derived from prostate-specific membrane
       antiqen (PSMA) and uses in treatment of prostate cancer)
     365470-51-5 365470-52-6 365470-53-7 365470-54-8 365470-55-9
IT
     365470-56-0
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (encoding PMSA peptide; immunogenic peptides derived from
       prostate-specific membrane antigen (PSMA) and uses in treatment of
       prostate cancer)
                  187968-05-4 187968-07-6 187968-08-7 187968-14-5
    187968-03-2
\operatorname{IT}
     187968-15-6
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); PRP (Properties); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (immunogenic peptides derived from prostate-specific membrane antigen
        (PSMA) and uses in treatment of prostate cancer)
     365490-33-1 3.65490-34-2 365490-35-3 365490-36-4
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IT
     365490-38-6
     RL: PRP (Properties)
        (unclaimed nucleotide sequence; immunogenic peptides derived from
       prostate-specific membrane antigen (PSMA) and uses thereof)
    187968-06-5
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     365424-41-5
     RL: PRP (Properties)
        (unclaimed sequence; immunogenic peptides derived from
       prostate-specific membrane antigen (PSMA) and uses thereof)
L85 ANSWER 4 OF 26 HCAPLUS COPYRIGHT 2003 ACS
     2001:598169 HCAPLUS
AN
     135:175408
DΝ
     Substances modulating FE65 interaction with hnRNPL and FEBPl for treatment
TI
     of neurodegenerative diseases
     Maury, Isabelle; Mercken, Luc; Fournier, Alain
ΙN
     Aventis Pharma S.A., Fr.
PA
     PCT Int. Appl., 51 pp.
SO
     CODEN: PIXXD2
DT
     Patent
     French
LΑ
     ICM C12N015-12
IC
         CO7KO14-47; C12Q001-68; C12N015-11; A61K038-00
     ICS
    1-11 (Pharmacology)
CC
     Section cross-reference(s): 3, 6
FAN.CNT 1
     PATENT NO.
                     KIND
                          DATE
                                          APPLICATION NO. DATE
    WO 2001059104 Al 20010816
                                     WO 2001-FR361 20010207
PΙ
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
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            LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
            DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                           20010817
                                                           20000210
                      A1
                                     FR 2000-1628
     FR 2804962
                                     BR 2001-8247
                           20021105
     BR 2001008247
                      Α
                                                           20010207
                                     EP 2001-907727 20010207
     EP 1257642
                           20021120
                      A1
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
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A1 20020523
                                           US 2001-780996
     US 2002061553
                                                            20010209
     NO 2002003783
                            20020909
                                           NO 2002-3783
                                                            20020809
                            20000210
PRAI FR 2000-1628
     US 2000-198500P P 20000418
                            20010207
     WO 2001-FR361
                       M
     Substances (peptides, nucleic acids, sugars,
AB
     lipids, antibodies) which modulate the interaction of amyloid precursor
     protein-binding protein FE65 with proteins hnRNPL and FEBPl and their use
     for treatment of neurodegenerative diseases are disclosed. Thus, using
     the yeast two hybrid system, fragments of hnRNPL and FEBP1 which bind to
     the PTB1 domain of FE65 were identified.
     hnRNPL FEBP1 fragment FE65 binding neurodegenerative disease treatment
ST
     Proteins, specific or class
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (FE65, PTB1 domain of; substances modulating FE65 interaction with
        hnRNPL and FEBP1 for treatment of neurodegenerative diseases)
    Antibodies
IT
     Carbohydrates, biological studies
     Lipids, biological studies
     Nucleic acids
     Peptides, biological studies
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (FE65-hnRNPL/FEBP1 interaction-modulating; substances modulating FE65
        interaction with hnRNPL and FEBP1 for treatment of neurodegenerative
        diseases)
     Proteins, specific or class
ΙT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (FEBP1; substances modulating FE65 interaction with hnRNPL and FEBP1
        for treatment of neurodegenerative diseases)
     Nervous system
IT
        (degeneration; substances modulating FE65 interaction with hnRNPL and
        FEBP1 for treatment of neurodegenerative diseases)
     cDNA sequences
IT
        (for FE65-binding fragments of human proteins hnRNPL and FEBP1)
     Proteins, specific or class
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (heterogeneous nuclear RNA-contg. ribonucleoprotein-assocd., hnRNPL;
        substances modulating FE65 interaction with hnRNPL and FEBP1 for
        treatment of neurodegenerative diseases)
    Genetic vectors
ΙT
    Virus vectors
        (hnRNPL/FEBP1 fragment-encoding; substances modulating FE65 interaction
        with hnRNPL and FEBP1 for treatment of neurodegenerative diseases)
     Protein sequences
IT
        (of FE65-binding fragments of human proteins hnRNPL and FEBP1)
     354643-19-9 354643-21-3
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); PRP
     (Properties); BIOL (Biological study); PROC (Process)
        (amino acid sequence; substances modulating FE65 interaction with
        hnRNPL and FEBP1 for treatment of neurodegenerative diseases)
     354643-18-8 354643-20-2
IT
     RL: BUU (Biological use, unclassified); PRP (Properties); BIOL (Biological
     study); USES (Uses)
        (nucleotide sequence; substances modulating FE65 interaction with
       hnRNPL and FEBP1 for treatment of neurodegenerative diseases)
     354645-01-5, 1: PN: WO0159104 SEQID: 1 unclaimed DNA 354645-03-7
IT
     354645-04-8 354645-05-9
     RL: PRP (Properties)
        (unclaimed nucleotide sequence; substances modulating FE65 interaction
        with hnRNPL and FEBP1 for treatment of neurodegenerative diseases)
```

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354645-02-6
IT
     RL: PRP (Properties)
        (unclaimed protein sequence; substances modulating FE65 interaction
        with hnRNPL and FEBP1 for treatment of neurodegenerative diseases)
              THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT 9
RE
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(2) Borg, J; MOLECULAR AND CELLULAR BIOLOGY 1996, V16/11, P6229
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(8) Zambrano, N; JOURNAL OF BIOLOGICAL CHEMISTRY 1997, V272/10, P6399
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L85 ANSWER 5 OF 26 HCAPLUS COPYRIGHT 2003 ACS
     2000:41751 HCAPLUS
AN
    132:304723
DN
     Influence of the type of junction in DNA-3'-peptide
\mathtt{TI}
     nucleic acid (PNA) chimeras on their binding
     affinity to DNA and RNA
     Greiner, Beate; Breipohl, Gerhard; Uhlmann, Eugen
ΑU
     Hoechst Marion Roussel Deutschland GmbH, Chemical Research G 838,
CS
     Frankfurt, D-65926, Germany
     Helvetica Chimica Acta (1999), 82(12), 2151-2159
SO
     CODEN: HCACAV; ISSN: 0018-019X
     Verlag Helvetica Chimica Acta
PB
     Journal
\mathsf{DT}
     English
LA
     6-2 (General Biochemistry)
     Section cross-reference(s): 33
     The automated online synthesis of a series of three DNA-3'-PNA (
ΑB
     PNA = Polyamide Nucleic Acids) chimeras is described, in which the
     3'-terminus of the oligonucleotide is linked to the amino terminus of the
     PNA via an N-(2-mercaptoethyl) - (X=S), N-(2-hydroxyethyl) - (X=O),
     or N-(2-aminoethyl)- (X=NH) N-[(thymin-1-yl)acetyl]glycine unit. In
     addn., the DNA-3'-PNA chimera with no nucleobase at the linking
     unit was prepd. The binding affinities of all chimeras were directly
     compared by detg. their Tm values in duplexes with complementary DNA, RNA,
     or DNA contg. a mismatch or abasic site opposite to the linker unit. We
     found that all chimeras in this study which have a nucleobase at the
     junction were able to form more stable duplexes with complementary DNA and
     RNA than the corresponding unmodified DNA. The influence of X on duplex
     stabilization was detd. to be 0 > S .apprxeq. NH, thus demonstrating the
     phosphodiester bridge to be the most favored linkage at the DNA/
     PNA junction. The strong duplex-destabilizing effects obsd. when
     base mismatches or non-basic sites were introduced opposite the nucleobase
     at the DNA/PNA junction, suggest that the base situated at the
     linking unit contributes significantly to duplex stabilization.
     peptide nucleic acid PNA binding
ST
     DNA RNA
     Molecular association
IT
        (binding affinity to complementary DNA and RNA sequences by DNA-3'-
        peptide nucleic acid (PNA)
        chimeras is influenced by nature of oligonucleotide-PNA
        junction)
     DNA
IT
     RNA
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
        (binding affinity to complementary DNA and RNA sequences by DNA-3'-
        peptide nucleic acid (PNA)
```

chimeras is influenced by nature of oligonucleotide-PNA

junction) Peptide nucleic acids ITRL: PEP (Physical, engineering or chemical process); SPN (Synthetic preparation); PREP (Preparation); PROC (Process) (binding affinity to complementary DNA and RNA sequences by DNA-3'peptide nucleic acid (PNA) chimeras is influenced by nature of oligonucleotide-PNA junction) Glass, biological studies IT RL: BPN (Biosynthetic preparation); BUU (Biological use, unclassified); BIOL (Biological study); PREP (Preparation); USES (Uses) (controlled pore, CPG; peptide nucleic acid conjugates; binding affinity to complementary DNA and RNA sequences by DNA-3'-peptide nucleic acid (PNA) chimeras is influenced by nature of oligonucleotide-PNA junction) 186050-42-0P 264893-89-2P 264893-91-6P 265075-78-3P IT RL: PEP (Physical, engineering or chemical process); SPN (Synthetic preparation); PREP (Preparation); PROC (Process) (binding affinity to complementary DNA and RNA sequences by DNA-3'peptide nucleic acid (PNA) chimeras is influenced by nature of oligonucleotide-PNA junction) 172405-23-1P TT RL: PNU (Preparation, unclassified); PREP (Preparation) (binding affinity to complementary DNA and RNA sequences by DNA-3'peptide nucleic acid (PNA) chimeras is influenced by nature of oligonucleotide-PNA junction) 105-36-2, Ethyl bromoacetate 141-43-5, reactions 156-57-0, IT 2-Mercaptoethylamine hydrochloride 563-96-2, Glyoxylic acid monohydrate 14470-28-1 20924-05-4 40615-36-9 82911-69-1 259827-32-2 RL: RCT (Reactant); RACT (Reactant or reagent) (binding affinity to complementary DNA and RNA sequences by DNA-3'peptide nucleic acid (PNA) chimeras is influenced by nature of oligonucleotide-PNA junction) 5835-28-9P, N-Hydroxyethyl glycine 141743-19-3P 172405-08-2P IT172405-33-3P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (binding affinity to complementary DNA and RNA sequences by DNA-3'peptide nucleic acid (PNA) chimeras is influenced by nature of oligonucleotide-PNA junction) THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD 13 RE.CNT RE (1) Bergmann, F; Tetrahedron Lett 1995, V36, P6823 HCAPLUS (2) Breipohl, G; Tetrahedron 1997, V53, P14671 (3) Egholm, M; Nature 1993, V365, P566 HCAPLUS (4) Hyrup, B; Bioorg Med Chem 1996, V4, P5 HCAPLUS (5) Matteucci, M; J Am Chem Soc 1981, V103, P3185 HCAPLUS (6) Nielsen, P; Science 1991, V254, P1497 HCAPLUS (7) Petersen, K; Bioorg Med Chem Lett 1995, V5, P1119 HCAPLUS (8) Uhlmann, E; Angew Chem, Int Ed 1996, V35, P2632 (9) Uhlmann, E; Angew Chem, Int Ed 1998, V37, P2796 HCAPLUS (10) Uhlmann, E; Biol Chem 1998, V379, P1045 HCAPLUS (11) Uhlmann, E; Peptide Nucleic Acids: Protocols and Applications 1999, P51 **HCAPLUS** (12) van der Laan, A; Bioorg Med Chem Lett 1998, V8, P663 HCAPLUS (13) Will, D; Tetrahedron 1995, V51, P12069 HCAPLUS

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1999:501533 HCAPLUS
AN
     132:194633
DN
TI
     PNA/DNA chimeras
     Uhlmann, Eugen; Greiner, Beate; Breipohl, Gerhard
ΑU
     Hoechst Marion Roussel Deutschland GmbH Chemical Research G 838, Frankfurt
CS
     am Main, D-65926, Germany
     Peptide Nucleic Acids (1999), 51-70. Editor(s): Nielsen, Peter E.;
SO
     Egholm, Michael. Publisher: Horizon Scientific Press, Norfolk, UK.
     CODEN: 67YLA6
     Conference
\mathsf{DT}
     English
LA
CC
     34-3 (Amino Acids, Peptides, and Proteins)
     Section cross-reference(s): 6, 33
     A convenient method for the solid-support synthesis of PNA/DNA
AB
     chimeras is described which makes use of monomethoxytrityl/acyl-protected
     monomeric building blocks. The acid-labile monomethoxytrityl (Mmt) group
     is employed for the temporary protection of the amino function of
     aminoethyl-glycine, while the exocyclic amino functions of the nucleobases
     are protected with ammonia-cleavable acyl protecting groups. This
     orthogonal protecting-group strategy is fully compatible with the std.
     phosphoramidite DNA synthesis method. The resulting PNA/DNA
     chimeras obey the Watson-Crick rules on binding to complementary DNA and
     RNA. Binding affinity of the PNA-DNA chimeras strongly depends
     on the PNA: DNA ratio. The PNA/DNA chimeras bind with
     higher affinity to RNA than to DNA, and the type of linking moiety between
     PNA and DNA could be adjusted to obtain optimal binding affinity.
     In addn. to their promising binding properties, PNA-DNA chimeras
     can also assume biol. functions, such as a primer function for DNA
     polymerases. Pure PNAs cannot induce RNase H cleavage of target
     RNA, which often supports the biol. efficacy of antisense agents.
     contrast, the DNA-PNA chimeras are able to stimulate cleavage of
     the target RNA by RNase H on formation of a RNA chimera duplex.
     PNA DNA chimera oligopeptide oligonucleotide prepn solid phase;
ST
     DNA PNA chimera oligopeptide oligonucleotide prepn solid phase;
     chimera PNA DNA oligopeptide oligonucleotide prepn solid phase;
     oligopeptide oligonucleotide PNA DNA chimera prepn solid phase;
     oligonucleotide oligopeptide chimera PNA DNA prepn solid phase
     Solid phase synthesis
IT
        (methods of prepn. of PNA/DNA chimeras using solid-phase
        synthesis)
     DNA
IT
     Oligonucleotides
       Peptide nucleic acids
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (methods of prepn. of PNA/DNA chimeras using solid-phase
        synthesis)
     Peptides, preparation
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (oligopeptides; methods of prepn. of PNA/DNA chimeras using
        solid-phase synthesis)
     259723-33-6P 259782-15-5P
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of as PNA/DNA chimeras using solid-phase synthesis)
              THERE ARE 37 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
        37
RĒ
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(2) Bergmann, F; Tetrahedron Lett 1995, V36, P6823 HCAPLUS
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(6) Breipohl, G; Innovation and Perspectives in Solid Phase Synthesis 1996, P61
    HCAPLUS
(7) Breipohl, G; Tetrahedron 1997, V53, P14671
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- L85 ANSWER 7 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 1999:310246 HCAPLUS
- DN 131:88176
- TI Synthesis of a monocharged **peptide nucleic**acid (PNA) analog and its recognition as substrate by
 DNA polymerases
- AU Lutz, M. J.; Will, D. W.; Breipohl, G.; Benner, S. A.; Uhlmann, E.
- CS Department of Chemistry, Swiss Federal Institute of Technology, Zurich, CH-8092, Switz.
- SO Nucleosides & Nucleotides (1999), 18(3), 393-401 CODEN: NUNUD5; ISSN: 0732-8311
- PB Marcel Dekker, Inc.
- DT Journal
- LA English
- CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 33
- AB The prepn. of a novel phosphoramidite monomer based on thyminyl acetic acid coupled to the secondary nitrogen of 2-(2-amino-ethyl-amino)ethanol is described. This monomer can be used to attach a deoxy-nucleotide to the carboxy terminus of a PNA oligomer by solid-phase synthesis. The resulting PNA primer is recognized as a substrate by various DNA polymerases.
- ST DNA transcription monocharged PNA primer; thyminyl acetic acid phosphoramidite prepn PNA DNA oligomer solidphase
- IT Avian myeloblastosis virus
 Coliphage T7
 Murine leukemia virus
 Pyrococcus furiosus
 Pyrococcus woesei

Thermus aquaticus

Thermus flavus Thermus thermophilus (recognition of monocharged peptide nucleic acid (PNA) analog substrate by DNA polymerase from) Escherichia coli IT(recognition of monocharged peptide nucleic acid (PNA) analog substrate by DNA polymerase from Klenow fragment of) DNA formation IT(replication; synthesis of a monocharged peptide nucleic acid (PNA) analog and its recognition as substrate by DNA polymerases) Reverse transcription ITSolid phase synthesis (synthesis of a monocharged peptide nucleic acid (PNA) analog and its recognition as substrate by DNA polymerases) Nucleic acids IT RL: MSC (Miscellaneous) (synthesis of a monocharged peptide nucleic acid (PNA) analog and its recognition as substrate by DNA polymerases) Peptide nucleic acids IT RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (synthesis of a monocharged peptide nucleic acid (PNA) analog and its recognition as substrate by DNA polymerases) 204692-17-1P 206435-20-3P · 229323-75-5P 204692-16-0P ITRL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (prepn. and reaction of in the synthesis of a monocharged peptide nucleic acid (PNA) analog for use as substrate by DNA polymerases) 229323-76-6P 229323-77-7P ITRL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (prepn. of for use as substrate of DNA polymerases for DNA transcription) 20924-05-4 74405-42-8D, solid-supported 89992-70-1 111-41-1 ITRL: RCT (Reactant); RACT (Reactant or reagent) (reaction of in the synthesis of a monocharged peptide nucleic acid (PNA) analog for use as substrate by DNA polymerases) 9012-90-2 9068-38-6 IT RL: CAT (Catalyst use); USES (Uses) (synthesis of a monocharged peptide nucleic acid (PNA) analog for use as substrate by) THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT RE (1) Breipohl, G; EP 0460446 HCAPLUS (2) Breipohl, G; Tetrahedron 1997, V53, P14671 (3) Egholm, M; J Am Chem Soc 1992, V114, P1895 HCAPLUS (4) Engels, J; DNA Sythesis in Biotechnology 1993, V2, P317 (5) Hyrup, B; Bioorg Med Chem 1996, V4, P5 HCAPLUS (6) Lutz, M; J Am Chem Soc 1997, V119, P3177 HCAPLUS (7) Nielsen, P; Science 1991, V254, P1497 HCAPLUS (8) Uhlmann, E; Angew Chem Int Ed Engl 1996, V108, P2793 (9) Uhlmann, E; Angew Chem Int Ed Engl 1998, V37, P2796 HCAPLUS (10) Uhlmann, E; Chem Rev 1990, V90, P543 HCAPLUS (11) Uhlmann, E; Nucleosides & Nucleotides 1997, V16, P603 HCAPLUS (12) Van der Laan, A; Bioorg Med Chem Lett 1998, V8, P663 HCAPLUS (13) Van der Laan, A; Tetrahedron Lett 1997, V38, P2249 HCAPLUS (14) Will, D; Tetrahedron 1995, V51, P12069 HCAPLUS

- L85 ANSWER 8 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 1999:91165 HCAPLUS
- TI Minimal modification of antisense oligonucleotides
- AU Uhlmann, E.
- CS Chemical Research, Hoechst Marion Roussel, Frankfurt, 65926, Germany
- SO Book of Abstracts, 217th ACS National Meeting, Anaheim, Calif., March 21-25 (1999), CARB-005 Publisher: American Chemical Society, Washington, D. C.
 - CODEN: 67GHA6
- DT Conference; Meeting Abstract
- LA English
- Uniformly phosphorothicate (PS) modified oligodeoxynucleotides (ODN) are AB antisense agents of the first generation. Although a no. of PS-ODN are in advanced stages of clin. development and the first antisense drug (Vitravene; Isis Pharmaceuticals) has been approved by the FDA, certain limitations of PS-ODN have emerged. Our approach to overcome these limitations is to reduce the no. of PS linkages within the ODN to a min. which is necessary to stabilize against nucleotlytic degrdn. We have developed a novel protection strategy which is a combination of the end-capping technique and the PS protection of internal pyrimidine positions which are the major sites of endonuclease degrdn. This protection scheme has successfully been used for specific inhibition of expression of various genes. Advantageously, it can also be combined with secondary modifications at the carbohydrate moieties, such as 2'-O-alkyl-modifications, or with partial replacement of the sugar phosphate backbone by 2-aminoethylglycine-based PNA units (peptide nucleic acid) leading to DNA-PNA chimeras.
- L85 ANSWER 9 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 1998:745539 HCAPLUS
- DN 130:66670
- TI PNA: synthetic polyamide nucleic acids with unusual binding properties
- AU Uhlmann, Eugen; Peyman, Anusch; Breipohl, Gerhard; Will, David W.
- CS Hoechst Marion Rouseel Deutschland GmbH, Frankfurt am Main, D-65926, Germany
- SO Angewandte Chemie, International Edition (1998), 37(20), 2796-2823 CODEN: ACIEF5; ISSN: 1433-7851
- PB Wiley-VCH Verlag GmbH
- DT Journal; General Review
- LA English
- CC 33-0 (Carbohydrates)
- A review with 160 refs. : since the investigation of oligonucleotides as AΒ potential therapeutics that target nucleic acids was initiated, the search for nucleic acid mimetics with improved properties, such as strengthened binding-affinity to complementary nucleic acids, increased biol. stability, and improved cellular uptake, has accelerated rapidly. Nielsen et al. first described what is undoubtedly one of the most interesting of the new derivs., the polyamide or peptide nucleic acids (PNAs), in which the entire sugar-phosphate backbone is replaced by an N-(2-aminoethyl)glycine polyamide structure. Since even minor structural changes in oligonucleotides, such as the replacement of an oxygen atom by sulfur (phosphorothioates), or by a neutral Me group (Me phosphonates), result in a decrease in binding affinity, it was even more astonishing to find that the drastic structural changes in PNAs result in nucleic acid mimetics with higher binding-affinity to complementary DNA and RNA than unmodified oligonucleotides. The remarkable binding properties of PNAs have spawned a rapidly expanding new field of research, where the targets are the synthesis of PNAs and PNA analogs, and their application as therapeutics, DNA diagnostics, and tools in

biotechnol. In add., investigation of **PNAs** and **PNA**/DNA chimeras can be used to generate information on the structural and biol. properties of DNA and RNA themselves. Furthermore, they may trigger the generation of new ideas on models for alternative living systems and potential transitions between different genetic systems.

ST review synthetic polyamide nucleic acid; polyamide nucleic acid review

IT DNA

RL: MSC (Miscellaneous); PNU (Preparation, unclassified); PREP (Preparation)

(PNA/DNA-chimeras; review of synthetic polyamide nucleic acids with unusual binding properties)

IT Nucleic acids

RL: MSC (Miscellaneous)

(review of synthetic polyamide nucleic acids with unusual binding properties)

IT Peptide nucleic acids

RL: MSC (Miscellaneous); PNU (Preparation, unclassified); PRP (Properties); PREP (Preparation)

(review of synthetic polyamide nucleic acids with unusual binding properties)

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L85
    ANSWER 10 OF 26 HCAPLUS COPYRIGHT 2003 ACS
     1998:667152 HCAPLUS
AN
     130:66764
DN
     DNA-PHONA-PNA chimeric molecules: contributions to binding
TI
     against complementary DNA
     Peyman, A.; Uhlmann, E.; Wagner, K.; Augustin, S.; Weiser, C.;
ΑU
     Hein, S.; Langner, D.; Breipohl, G.; Will, D. W.
     Hoechst Marion Roussel Deutschland GmbH, Frankfurt, D-65926, Germany
CS
     Nucleosides & Nucleotides (1998), 17(9-11), 1997-2001
SO
     CODEN: NUNUD5; ISSN: 0732-8311
     Marcel Dekker, Inc.
PΒ
     Journal
\operatorname{DT}
     English
LA
     34-3 (Amino Acids, Peptides, and Proteins)
CC
     Section cross-reference(s): 33
     The synthesis of a DNA-phosphonate peptide nucleic
AB
     acid analog (PHONA) -peptide nucleic
     acid (PNA) chimeric mol. using a monomethoxytrityl (Mmt)
     protection strategy is described. The chimeric oligomer shows duplex
     binding properties that are comparable to the corresponding PNA.
     Thus, PHONA building blocks can be incorporated into PNAs
     without distortion of the PNA structure.
     peptide nucleic acid phosphonate analog
ST
     prepn DNA binding
ΙT
     DNA
     RL: PRP (Properties)
        (complexes, with peptide nucleic acid-
        peptide nucleic acid phosphonate analogs;
        prepn. and DNA binding properties of DNA-peptide
        nucleic acid phosphonate analog-peptide
        nucleic acid chimeric mols.)
     Peptide nucleic acids
ΙT
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (phosphonate backbone analogs; prepn. and DNA binding properties of
        DNA-peptide nucleic acid phosphonate
        analog-peptide nucleic acid chimeric
        mols.)
     217636-83-4P 217636-84-5P
                                  217636-85-6P 217636-86-7P
\operatorname{IT}
                                                                 217636-87-8P
     217636-88-9P 217636-89-0P
                                   217636-90-3P 217636-91-4P
                                                                 217636-92-5P
     217636-93-6P 217636-94-7P
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and DNA binding properties of DNA-peptide
        nucleic acid phosphonate analog-peptide
        nucleic acid chimeric mols.)
     217636-80-1P 217636-81-2P 217636-82-3P
IT
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (prepn. and DNA binding properties of DNA-peptide
        nucleic acid phosphonate analog-peptide
        nucleic acid chimeric mols.)
RE.CNT
              THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD
        10
RE
(1) de Mesmaker, A; Acc Chem Res 1995, V28, P366
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- (6) Uhlmann, E; Angew Chem Int Ed Engl 1996, V35, P2632
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- ANSWER 11 OF 26 HCAPLUS COPYRIGHT 2003 ACS L85
- 1998:618936 HCAPLUS AN
- 129:227036 DN
- Peptide nucleic acids (PNA) and TI PNA-DNA chimeras. From high binding affinity towards biological function
- Uhlmann, Eugen ΑU
- CS Hoechst Marion Roussel Deutschland G.m.b.H., Frankfurt/Main, D-65926, Germany
- Biological Chemistry (1998), 379(8/9), 1045-1052 SO CODEN: BICHF3; ISSN: 1431-6730
- Walter de Gruyter & Co. PB
- Journal; General Review DT
- English LA
- 6-0 (General Biochemistry) CC
- A review is given with 45 refs. Oligonucleotide analogs are of major AΒ interest as tools in mol. biol., as diagnostics, and as potential pharmaceuticals which bind in a predictable way to certain nucleic acid target sequences, aiming at the inhibition of expression of disease-causing genes. One of the most promising nucleic acid mimetics are the peptide- or polyamide- nucleic acids (PNA) which bind with higher affinity to DNA and RNA than natural oligonucleotides. In these non-ionic PNAs, the entire sugar-phosphate backbone is replaced by an N-amino-ethylglycine-based polyamide structure. A unique property of PNA is its ability to displace one strand of a DNA double-helix. This strand displacement process, which is inefficient with DNA, is supported by the formation of an unusually stable internal (PNA), DNA triple helix. The combination of PNA and DNA in 1 mol. results in PNA/DNA chimeras with new properties. They show improved aq. soly. compared to pure PNAs due to their partially neg. charged structure. The cellular uptake of the chimeras is better than of pure PNAs. In contrast to PNA, the chimeras bind exclusively in the antiparallel orientation under physiol. conditions. The binding affinity is generally stronger when the PNA/DNA chimeras are hybridized to RNA than to DNA, whereby the strength of binding strongly depends on the PNA: DNA ratio. PNA/DNA chimeras are recognized as substrates by various nucleic acid processing enzymes, and consequently can also assume biol. functions, such as a primer function for DNA polymerases. Pure PNA cannot induce RNase H cleavage of target RNA, which is believed to support the biol. efficacy of antisense agents. DNA-PNA chimeras are able to stimulate cleavage of the target RNA by RNase H upon formation of an RNA chimera duplex.
- review peptide nucleic acid DNA chimera ST
- Antisense oligonucleotides ITDNA

Peptide nucleic acids

RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process)

(peptide nucleic acids and PNA

-DNA chimeras)

9012-90-2, DNA polymerase IT

RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)

(peptide nucleic acids and PNA -DNA chimeras)

DN 128:321903

TI Optimization of the binding properties of PNA-(5')-DNA chimerae

AU van der Laan, A. C.; Havenaar, P.; Oosting, R. S.; Kuyl-Yeheskiely, E.; Uhlmann, E.; van Boom, J. H.

CS Gorlaeus Lab., Leiden Inst. of Chemistry, Leiden, 2300 RA, Neth.

SO Bioorganic & Medicinal Chemistry Letters (1998), 8(6), 663-668 CODEN: BMCLE8; ISSN: 0960-894X

PB Elsevier Science Ltd.

DT Journal

LA English

CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 3, 33

GI

AB The synthesis and evaluation of PNA-(5')-DNA chimera contg.
either a 5'-amide (i.e. I; T = thymin-1-yl), a 5'-phosphodiester (i.e. II)
or 5'-phosphonate linkages (i.e. III; R = H, thymin-1-ylacetyl) at the
junction site are described. The 5'-linkages were installed using
protected phosphoramidite and phosphonate building blocks. It is shown
that PNA-(5')-DNA of types I, II, and III (R =
thymin-1-ylacetyl) have a higher binding affinity with complementary RNA
than native DNA, and that the antisense activity is mainly due to RNase H.
ST peptide nucleic acid DNA chimera prepn;

ST peptide nucleic acid DNA chimera prepn; binding property optimization PNA DNA; structure activity PNA DNA binding

IT Structure-activity relationship

(DNA-binding; prepn. and optimization of PNA-DNA chimera binding properties)

IT DNA

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (complexes; prepn. and optimization of PNA-DNA chimera binding properties)

IT Translation, genetic

(prepn. and optimization of PNA-DNA chimera binding

```
properties)
    Antisense DNA
\operatorname{IT}
    RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); BIOL (Biological
     study); PREP (Preparation)
        (prepn. and optimization of PNA-DNA chimera binding
       properties)
    Peptide nucleic acids
ΙT
    RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and optimization of PNA-DNA chimera binding
       properties)
                   207020-61-9P 207020-62-0P 207020-63-1P 207020-64-2P
    207020-60-8P
IT
    207020-65-3P
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); SPN (Synthetic preparation); BIOL (Biological
     study); PREP (Preparation)
        (prepn. and optimization of PNA-DNA chimera binding
       properties)
    9050-76-4
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (prepn. and optimization of PNA-DNA chimera binding
       properties)
                                              207020-38-0
                                                            207020-46-0
    207020-32-4 207020-33-5 207020-37-9
IT
     207020-47-1 207020-51-7 207020-52-8
     RL: PRP (Properties)
        (prepn. and optimization of PNA-DNA chimera binding
       properties)
    207020-34-6P 207020-35-7P 207020-36-8P 207020-39-1P
                                                               207020-40-4P
IT
     207020-41-5P 207020-42-6P
                                  207020-43-7P
                                                 207020-44-8P
                                                                207020-45-9P
     207020-48-2P 207020-49-3P 207020-50-6P
                                                                207020-54-0P
                                                 207020-53-9P
    207020-55-1P 207020-56-2P 207020-57-3P 207020-58-4P 207020-59-5P
     RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
        (prepn. and optimization of PNA-DNA chimera binding
       properties)
     4712-55-4 10242-36-1 20924-05-4 57260-73-8 172316-36-8
ΙT
                                              206435-20-3
    172316-42-6 172316-44-8 182998-85-2
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. and optimization of PNA-DNA chimera binding
       properties)
                                                               206435-23-6P
                                 206435-21-4P
                                                 206435-22-5P
    203643-20-3P 203643-39-4P
IT
    206435-24-7P 206435-25-8P 206435-26-9P 206435-27-0P
                                                               206435-28-1P
     206435-29-2P 206887-50-5P 206887-51-6P 206887-52-7P 206887-53-8P
     206887-54-9P 206887-55-0P 207020-28-8P 207020-29-9P 207020-30-2P
     207020-31-3P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (prepn. and optimization of PNA-DNA chimera binding
       properties)
    ANSWER 13 OF 26 HCAPLUS COPYRIGHT 2003 ACS
L85
    1998:186571 HCAPLUS
AN
    128:240314
DN
     A nucleic acid amplification method using peptide
TI
     nucleic acids as primers for thermostable DNA
     polymerases
     Uhlmann, Eugen; Breipohl, Gerhard; Benner, Steven;
IN
     Lutz, Michael
     Hoechst A.-G., Germany
PA
SO
     Eur. Pat. Appl., 17 pp.
     CODEN: EPXXDW
     Patent
\mathsf{DT}
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German

LA

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ICM C12Q001-68
IC
    3-1 (Biochemical Genetics)
CC
FAN.CNT 1
    PATENT NO. KIND DATE
                                          APPLICATION NO. DATE
    EP 829542 A1 19980318 EP 1997-115521 19970908
PI
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, FI
    DE 19637339
                     A1 19980319
                                          DE 1996-19637339 19960913
    US 6063571 A 20000516
CA 2215489 AA 19980313
JP 10099088 A2 19980421
                                          US 1997-927274 19970911
                                         -CA 1997-2215489 19970912
                                          JP 1997-250443 19970916
PRAI DE 1996-19637339
                           19960913
    A method of using peptide nucleic acids (
AB
    PNAs) as primers for DNA amplification with thermostable DNA
    polymerases, i.e. in PCR, is described. The only modification to the
    PNAs that is essential is the introduction of 1-3 3'-terminal
    deoxynucleotides with a free 3'-hydroxyl group. Methods for the synthesis
    of deoxynucleotide-terminated primers are also given.
    peptide nucleic acid primer PCR; PNA
ST
    deoxynucleotide terminated PCR primer
    Peptide nucleic acids
IT
    RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses)
        (3'-ologodeoxynucleotide with 3'-OH-contg., as primers; nucleic acid
        amplification method using peptide nucleic
        acids as primers for thermostable DNA polymerases)
    PCR (polymerase chain reaction)
TT
        (nucleic acid amplification method using peptide
       nucleic acids as primers for thermostable DNA
       polymerases)
    204692-16-0P 204692-17-1P 204692-18-2P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (prepn. and reactions of, in prepn. peptide nucleic
       acids; nucleic acid amplification method using peptide
       nucleic acids as primers for thermostable DNA
       polymerases)
    204867-94-7P 204867-95-8P 204867-96-9P
IT
    RL: ARG (Analytical reagent use); SPN (Synthetic preparation); ANST
     (Analytical study); PREP (Preparation); USES (Uses)
        (prepn. of, as PCR primer; nucleic acid amplification method using
       peptide nucleic acids as primers for
       thermostable DNA polymerases)
               7087-68-5P, Diisopropylethylamine 14470-28-1P 20924-05-4P
    111-41-1P
IT
    89992-70-1P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (reactions of, in prepn. peptide nucleic
       acids; nucleic acid amplification method using peptide
       nucleic acids as primers for thermostable DNA
       polymerases)
     9012-90-2, DNA polymerase
IT
    RL: ARG (Analytical reagent use); CAT (Catalyst use); ANST (Analytical
     study); USES (Uses)
        (thermostable, peptide nucleic acid-based
        primers for; nucleic acid amplification method using peptide
       nucleic acids as primers for thermostable DNA
       polymerases)
L85 ANSWER 14 OF 26 HCAPLUS COPYRIGHT 2003 ACS
    1998:70167 HCAPLUS
AN
    128:167687
DN
```

PHONA - PNA co-oligomers: nucleic acid mimetics with interesting

TI

properties

AU Peyman, Anusch; Uhlmann, Eugen; Wagner, Konrad; Augustin, Sascha; Weiser, Caroline; Will, David W.; Breipohl, Gerhard

CS Hoechst Marion Roussel Deutschland GmbH, Frankfurt, D-65926, Germany

Angewandte Chemie, International Edition in English (1998), Volume Date 1997, 36(24), 2809-2812

CODEN: ACIEAY; ISSN: 0570-0833

PB Wiley-VCH Verlag GmbH

DT Journal

LA English

CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 6, 33

GI

HO (CH₂)
$$6$$
NH $\frac{O}{O}$ $\frac{O}{O}$ $\frac{O}{O}$ $\frac{N}{H}$ $\frac{Ac}{6}$ $\frac{I}{I}$

AB Alternating title co-oligomer I contg. peptide nucleic acid (PNA) and (aminomethyl)phosphonic acid backbones was prepd. and melting temps. (Tm) of complexes with completely or partially complementary DNA measured. The binding properties of I with complementary DNA are very similar to those of PNAs, but the co-oligomer I has a much better water soly.

ST aminomethylphosphonate peptide nucleic acid prepn stability; DNA complex aminomethylphosphonate peptide nucleic acid

IT Peptide nucleic acids

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (prepn. of (aminomethyl)phosphonic acid backbone peptide nucleic acid co-oligomers as nucleic acid mimetics with interesting properties)

IT 20924-05-4, 1-Thyminylacetic acid 57260-73-8, N-tert-Butoxycarbonylethylenediamine 85363-76-4 172405-31-1 RL: RCT (Reactant); RACT (Reactant or reagent)

(prepn. of (aminomethyl)phosphonic acid backbone peptide nucleic acid co-oligomers as nucleic acid mimetics with interesting properties)

IT 185670-76-2P 185670-78-4P 185670-79-5P 202914-62-3P 202914-63-4P 202914-64-5P 202914-65-6P 202914-66-7P 202914-67-8P 202914-68-9P 202914-69-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. of (aminomethyl)phosphonic acid backbone peptide nucleic acid co-oligomers as nucleic acid mimetics with interesting properties)

RE.CNT 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD RE

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- (33) Will, D; Tetrahedron 1995, V51, P12069 HCAPLUS
- L85 ANSWER 15 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 1997:758327 HCAPLUS
 - Correction of: 1997:714702
- DN 127:346655
 - Correction of: 127:319261
- TI Novel synthetic routes to **PNA** monomers and **PNA-**DNA linker molecules
- AU Breipohl, Gerhard; Will, David W.; Peyman, Anusch; Uhlmann, Eugen
- CS Hoechst Marion Roussel Deutschland GmbH, Frankfurt am Main, D-65926, Germany
- SO Tetrahedron (1997), 53(43), 14671-14686 CODEN: TETRAB; ISSN: 0040-4020
- PB Elsevier
- DT Journal
- LA English
- CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 33

GI

- AB Novel methods for the prepn. of monomethoxytrityl (Mmt)-protected aminoethylglycine building blocks [I; B = 1-thyminyl, N4-(4-methoxybenzoyl)-1-cytosinyl, N6-(4-methoxybenzoyl)-9-adeninyl, N2-acetyl-06-diphenylcarbamoyl-9-guaninyl, N2-isobutyryl-9-guaninyl) and dimethoxytrityl (Dmt)-protected hydroxyethylglycine derivs. II, useful for the synthesis of polyamide nucleic acids (PNAs) and PNA /DNA chimeras, are described. The protecting group strategy employed for PNA monomer synthesis produces intermediates that are easily isolated, minimizes chromatog, purifn., and is suitable for large-scale monomer synthesis.
- ST peptide nucleic acid monomer prepn; nucleic acid polyamide protected building block
- IT Peptide nucleic acids

RL: SPN (Synthetic preparation); PREP (Preparation) (novel synthetic routes to protected PNA monomers and PNA-DNA linker mols.)

IT 96-32-2 107-15-3, 1,2-Ethanediamine, reactions 107-59-5 141-43-5,
reactions 2916-14-5 3891-07-4, N-(2-Hydroxyethyl)phthalimide
5292-43-3 20924-05-4 112233-74-6 172405-10-6 172405-18-4
172405-20-8

RL: RCT (Reactant); RACT (Reactant or reagent)
(novel synthetic routes to protected PNA monomers and
PNA-DNA linker mols.)

 IT
 66937-71-1P
 153765-10-7P
 172405-24-2P
 172405-25-3P
 172405-32-2P

 172729-41-8P
 184241-26-7P
 188779-49-9P
 188779-50-2P
 188779-51-3P

 188779-53-5P
 188779-54-6P
 188779-56-8P
 188779-57-9P
 188779-58-0P

 188779-59-1P
 188779-60-4P
 188779-61-5P
 188779-62-6P
 188779-63-7P

 197801-81-3P
 197801-83-5P
 197801-98-2P
 197802-00-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(novel synthetic routes to protected **PNA** monomers and **PNA-DNA** linker mols.)

IT 170490-73-0P 172316-36-8P 172316-40-4P 172316-41-5P 172316-42-6P 185810-72-4P 185810-73-5P 185810-74-6P 188779-64-8P RL: SPN (Synthetic preparation); PREP (Preparation)

(novel synthetic routes to protected PNA monomers and PNA-DNA linker mols.)

- L85 ANSWER 16 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 1997:714702 HCAPLUS
- DN 127:319261
- TI Novel synthetic routes to PNA monomers and PNA-DNA linker molecules
- AU Breipohl, Gerhard; Will, David W.; Peyman, Anusch; Uhimann, Eugen
- CS Hoechst Marion Roussel Deutschland GmbH, Frankfurt am Main, D-65926, Germany
- SO Tetrahedron (1997), 53(43), 14671-14686 CODEN: TETRAB; ISSN: 0040-4020
- PB Elsevier
- DT Journal
- LA English

CC 34-3 (Amino Acids, Peptides, and Proteins) Section cross-reference(s): 26

GI

- Novel methods for the prepn. of monomethoxytrityl (Mmt)-protected aminoethylglycine building blocks I [B = 1-thyminyl, N4-(4-methoxybenzoyl)-1-cytosinyl, N6-(4-methoxybenzoyl)-9-adeninyl, N2-acetyl-O6-diphenylcarbamoyl-9-guaninyl, N2-isobutyryl-9-guaninyl] and dimethoxytrityl (Dmt)-protected hydroxyethylglycine derivs. II, useful for the synthesis of polyamide nucleic acids (PNAs) and PNA/DNA chimeras are described. The protecting group strategy employed for PNA monomer synthesis produces easily isolable intermediates, minimizes chromatog. purifn., and is suitable for large-scale monomer synthesis.
- ST peptide nucleic acid monomer prepn; protected building block polyamide nucleic acid
- IT Peptide nucleic acids

RL: SPN (Synthetic preparation); PREP (Preparation) (novel synthetic routes to protected PNA monomers and PNA-DNA linker mols.)

IT 96-32-2, Methyl bromoacetate 107-15-3, 1,2-Ethanediamine, reactions
107-59-5, tert-Butyl chloroacetate 141-43-5, reactions 2916-14-5,
Allyl chloroacetate 3891-07-4, N-(2-Hydroxyethyl)phthalimide
5292-43-3, tert-Butyl bromoacetate 20924-05-4 112233-74-6
172405-10-6 172405-18-4 172405-20-8
RL: RCT (Reactant); RACT (Reactant or reagent)

(novel synthetic routes to protected PNA monomers and PNA-DNA linker mols.)

IT 66937-71-1P 153765-10-7P 172405-24-2P 172405-25-3P 172405-32-2P 172729-41-8P 184241-26-7P 188779-49-9P 188779-50-2P 188779-51-3P 188779-53-5P 188779-54-6P 188779-56-8P 188779-57-9P 188779-58-0P 188779-59-1P 188779-60-4P 188779-61-5P 188779-62-6P 188779-63-7P 197801-81-3P 197801-83-5P 197801-98-2P 197802-00-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(novel synthetic routes to protected PNA monomers and PNA-DNA linker mols.)

(novel synthetic routes to protected **PNA** monomers and **PNA-**DNA linker mols.)

- L85 ANSWER 17 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 1997:591221 HCAPLUS
- DN 127:262910
- TI Synthesis of polyamide nucleic acids (PNAs), PNA /DNA-chimeras and phosphonic ester nucleic acids (PHONAs)
- AU Uhlmann, E.; Will, D. W.; Breipohl, G.; Peyman, A.; Langner, D.; Knolle, J.; O'Malley, G.
- CS Central Pharma Res., Hoechst AG, Frankfurt, D-65926, Germany
- SO Nucleosides & Nucleotides (1997), 16(5 & 6), 603-608

```
CODEN: NUNUD5; ISSN: 0732-8311
     Dekker
PB
     Journal; General Review
DΤ
     English
LA
     33-0 (Carbohydrates)
CC
     Section cross-reference(s): 34
     A review with 18 refs. on methods for the prepn. of polyamide nucleic
AB
     acids (PNAs) and derivs. thereof by different synthetic routes
     is described. The first strategy makes use of 9-Fluorenylmethoxycarbonyl
     (Fmoc)/monomethoxytrityl (Mmt) protected building blocks, whereas the
     second approach involves the use of Mmt/acyl protected monomers, which
     allows the prepn. of PNA/DNA chimera. Addnl., a block coupling
     strategy is presented for the synthesis of novel phosphonic ester nucleic
     acids (PHONAs).
ST
     monomethoxytrityl protective group DNA prepn review;
     fluorenylmethoxycarbonyl protective group DNA prepn review; phosphonic
     ester nucleic acid prepn review; PNA DNA chimera prepn review;
     polyamide nucleic acid DNA chimera review
     Protective groups
IT
        (Fmoc/MMTr; prepn. of polyamide nucleic acids, PNA
        /DNA-chimeras and phosphonic ester nucleic acids)
     Peptide nucleic acids
ΙT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (PNA/DNA-chimeras; prepn. of polyamide nucleic acids,
        PNA/DNA-chimeras and phosphonic ester nucleic acids)
ΙT
     DNA
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (PNA/DNA-chimeras; prepn. of polyamide nucleic acids,
        PNA/DNA-chimeras, and phosphonic ester nucleic acids)
     Nucleic acids
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (phosphonic ester; prepn. of polyamide nucleic acids, PNA
        /DNA-chimeras, and phosphonic ester nucleic acids)
    ANSWER 18 OF 26 HCAPLUS COPYRIGHT 2003 ACS
L85
     1997:412349 HCAPLUS
AN
DN
     127:66087
     Solid-phase synthesis of PNA-DNA chimeric oligomers
TI
     Will, D.W.; Breipohl, G.; Langner, D.; Uhlmann,
ΑU
CS
     Hoechst AG, Allgemeine Pharma Forschung G838, Frankfurt am Main, D-65926,
     Germany
SO
     Innovation and Perspectives in Solid Phase Synthesis & Combinatorial
     Libraries: Peptides, Proteins and Nucleic Acids -- Small Molecule Organic
     Chemical Diversity, Collected Papers, International Symposium, 4th,
     Edinburgh, Sept. 12-16, 1995 (1996), Meeting Date 1995, 65-68. Editor(s):
     Epton, Roger. Publisher: Mayflower Scientific, Birmingham, UK.
     CODEN: 640NA9
     Conference
DT
     English
LA
     33-10 (Carbohydrates)
CC
     Section cross-reference(s): 34
     A symposium on PNA-DNA chimeric oligomers have been prepd. using
AΒ
     automated solid-phase prepn. A novel Mmt protecting-group strategy for
     the PNA part of the mol. was employed which allowed the use of
     std. DNA synthesis and deprotection chem.
     monomethoxytrityl protecting group DNA PNA symposium;
ST
     PNA DNA solid phase prepn symposium
     Protective groups
IT
        (monomethoxytrityl; solid phase prepn. of PNA/DNA chimeric
        oligomers)
     Solid phase synthesis
IT
        (solid phase prepn. of PNA/DNA chimeric oligomers)
```

IT DNA

Peptide nucleic acids

RL: SPN (Synthetic preparation); PREP (Preparation) (solid phase prepn. of PNA/DNA chimeric oligomers)

- L85 ANSWER 19 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 1997:412348 HCAPLUS
- DN 127:66086
- TI Synthesis of polyamide nucleic acids using a new protection scheme which is fully compatible with oligonucleotide synthesis
- AU Breipohl, G.; Will, D.W.; Langner, D.; Knolle, J.; Uhlmann, E.
- CS Hoechst AG, Allgemeine Pharma Forschung G838, Frankfurt am Main, D-65926, Germany
- Innovation and Perspectives in Solid Phase Synthesis & Combinatorial Libraries: Peptides, Proteins and Nucleic Acids--Small Molecule Organic Chemical Diversity, Collected Papers, International Symposium, 4th, Edinburgh, Sept. 12-16, 1995 (1996), Meeting Date 1995, 61-64. Editor(s): Epton, Roger. Publisher: Mayflower Scientific, Birmingham, UK. CODEN: 640NA9
- DT Conference
- LA English
- CC 33-10 (Carbohydrates)
- AB A symposium on the prepn. of novel monomethoxytrityl (Mmt) protected monomers for the prepn. of polyamide nucleic acids (PNAs) is described. Use of the acid-labile Mmt group as temporary protection for the primary amino function of aminoethylglycine in combination with base-labile acyl-type protecting groups for the nucleobases allow a synthetic strategy similar to std. oligo-nucleotide synthesis conditions. PNAs of mixed base sequence have been synthesized with this method.
- ST monomethoxytrityl protective group nucleic acid symposium; polyamide nucleic acid prepn symposium
- IT Protective groups

(monomethoxytrityl; prepn. of polyamide nucleic acids using a new protection which is fully compatible with oligodeoxyribonucleotide prepn.)

IT Nucleic acids

RL: SPN (Synthetic preparation); PREP (Preparation) (polyamide; prepn. of polyamide nucleic acids using a new protection which is fully compatible with oligodeoxyribonucleotide prepn.)

- L85 ANSWER 20 OF 26 HCAPLUS COPYRIGHT 2003 ACS
- AN 1997:283607 HCAPLUS
- DN 126:264359
- TI Preparation of ethylglycine derivatives
- IN Breipohl, Gerhard; Uhlmann, Eugen; Will, David
 William
- PA Hoechst A.-G., Germany
- SO Ger. Offen., 14 pp.

CODEN: GWXXBX

- DT Patent
- LA German
- IC ICM C07K005-078
- CC 34-3 (Amino Acids, Peptides, and Proteins)

Section cross-reference(s): 6

PAN.	CNT I			•	
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
ΡI	DE 19532553	A1	19970306	DE 1995-19532553	19950904
	EP 761681	A2	19970312	EP 1996-113530	19960822
	EP 761681	А3	19970709		
	EP 761681	B1	20020313		

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R: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
                           20020315
                                         AT 1996-113530
                                                          19960822
    AT 214398
                      T3 20021016
                                          ES 1996-113530 19960822
    ES 2173230
                      Al 19970306
                                                         19960902
    AU 9664408
                                         AU 1996-64408
                      B2 19990729
    AU 708034
                                          CA 1996-2184681 19960903
    CA 2184681
                      AA 19970305
                                          NO 1996-3677 19960903
    NO 9603677
                          19970305
                      A
                                          JP 1996-232692 19960903
                      A2 19970513
    JP 09124572
    US 5817811
                          19981006
                                          US 1996-707149
                                                         19960903
                      A
PRAI DE 1995-19532553 A
                           19950904
OS
    MARPAT 126:264359
    N-ethylglycine derivs. PG-X-CH2CH2N(COCH2B1)CH2CO2H (PG is a urethane- or
AB
    trityl-type amino protecting group which is cleavable by weak acid; X = NH
    or O; B1 = nucleotide base in which exocyclic amino or hydroxy groups are
    protected), useful in PNA or PNA/DNA hybrid prepn.,
    were prepd. Thus, 2-aminoethanol was condensed with bromoacetic acid t-Bu
    ester, then with thyminylacetic acid, the product deesterified, and the
    acid treated with DMT-Cl to give a protected PNA monomer.
    ethylglycine prepn; glycine ethyl prepn; aminoethanol condensation
ŞΤ
    bromoacetate thyminylacetic acid; PNA DNA hybrid prepn
    Peptide nucleic acids
IT
    RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
     (Preparation)
        (precursor prepn; prepn of ethylglycine derivs useful in PNA
       or PNA/DNA hybrid synthesis)
    Condensation reaction
IT
        (prepn of ethylglycine derivs useful in PNA or PNA
       /DNA hybrid synthesis)
    66937-71-1P 104732-23-2P 172405-25-3P 172405-32-2P
                                                              172729-41-8P
IT
    188779-49-9P 188779-50-2P 188779-51-3P 188779-53-5P
                                                              188779-55-7P
    188779-56-8P 188779-57-9P 188779-58-0P 188779-59-1P 188779-60-4P
    188779-61-5P 188779-63-7P
    RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic
    preparation); PREP (Preparation); RACT (Reactant or reagent)
        (prepn of ethylglycine derivs useful in PNA or PNA
       /DNA hybrid synthesis)
    170490-73-0P 172316-36-8P
                                  172316-40-4P 172316-41-5P 185810-72-4P
\operatorname{IT}
    185810-73-5P 188779-64-8P
    RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
     (Preparation)
        (prepn of ethylglycine derivs useful in PNA or PNA
       /DNA hybrid synthesis)
    107-15-3, 1,2-Ethanediamine, reactions 107-59-5, Chloroacetic acid,
IT
    tert-butyl ester 141-43-5, reactions 5292-43-3, Bromoacetic acid,
    tert-butyl ester 20924-05-4 172405-10-6 172405-18-4
                                                              188779-52-4
    188779-62-6
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn of ethylglycine derivs useful in PNA or PNA
       /DNA hybrid synthesis)
    ANSWER 21 OF 26 HCAPLUS COPYRIGHT 2003 ACS
L85
    1997:224058 HCAPLUS
AN
    126:274010
DN
    Recognition of Uncharged Polyamide-Linked Nucleic Acid Analogs by DNA
TI
    Polymerases and Reverse Transcriptases
    Lutz, Michael J.; Benner, Steven A.; Hein, Silvia; Breipohl,
ΑU
    Gerhard; Uhlmann, Eugen
     Department of Chemistry, Swiss Federal Institute of Technology, Zurich,
CS
```

Journal of the American Chemical Society (1997), 119(13), 3177-3178

PB American Chemical Society
DT Journal

SO

CH-8092, Switz.

CODEN: JACSAT; ISSN: 0002-7863

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English
LA
    7-3 (Enzymes)
CC
     Polyamide-linked nucleic acid (PNAs) are DNA mimics in which the
AB
     deoxyribose phosphate backbone is replaced by uncharged
     N-(2-aminoethyl) glycine units. Here, the authors report that several DNA
     polymerases and reverse transcriptases are able to elongate a PNA
     primer with a nucleophilic 3'-hydroxyl group, despite the fact that no
     phosphate residues are present in the PNA primer to interact
     with the polymerase. Enzymic synthesis of PNA-DNA chimeras
    might have implications for the use of modified PNAs in advanced
     diagnostic systems, allowing facilitated screening for genetic mutations,
     and as tools for studying structure-function relationships in enzymes that
     process nucleic acids. These results are also interesting in the light of
    models for the origin of life that propose an evolutionary linkage between
     a PNA-like and a DNA-protein world.
    peptide nucleic acid primer polymerase
    transcriptase; DNA polymerase primer peptide nucleic
     acid; reverse transcriptase primer peptide
     nucleic acid
     Reverse transcription
ΙŢ
        (recognition of uncharged DNA mimics (peptide nucleic
        acid primers) by DNA polymerases and reverse transcriptases)
    Peptide nucleic acids
ΙT
     Primers (nucleic acid)
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (recognition of uncharged DNA mimics (peptide nucleic
        acid primers) by DNA polymerases and reverse transcriptases)
     DNA formation
IT
        (replication; recognition of uncharged DNA mimics (peptide
       nucleic acid primers) by DNA polymerases and reverse
       transcriptases)
     9012-90-2, DNA polymerase
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (I; recognition of uncharged DNA mimics (peptide
       nucleic acid primers) by DNA polymerases and reverse
        transcriptases)
     9068-38-6, Reverse transcriptase 188901-47-5
IT
     RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (recognition of uncharged DNA mimics (peptide nucleic
        acid primers) by DNA polymerases and reverse transcriptases)
L85 ANSWER 22 OF 26 HCAPLUS COPYRIGHT 2003 ACS
    1997:88503 HCAPLUS
AN
    126:100903
DN
    Phosphonomonoester nucleic acids, process for their preparation, and their
TI
    use in molecular biology and as pharmaceuticals
     Peyman, Anuschirwan; Uhlmann, Eugen; Breipohl, Gerhard
IN
     ; Wallmeier, Holger
    Hoechst A.-G., Germany
PA
    Can. Pat. Appl., 126 pp.
SO
    CODEN: CPXXEB
    Patent
DT
LA
    English
IC
    ICM C12Q001-68
     ICS C07K002-00; C07H021-00; A61K048-00; A61K031-70; A61K038-00
     6-2 (General Biochemistry)
CC
     Section cross-reference(s): 1, 3, 33
FAN.CNT 2
     PATENT NO. KIND DATE
                                          APPLICATION NO. DATE
```

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CA 2171589
                      AA 19960914
                                           CA 1996-2171589 19960312
PI
                       A1 19960919
                                           DE 1995-19508923 19950313
     DE 19508923
     DE 19543865
                       A1 19970605
                                            DE 1995-19543865 19951124
PRAI DE 1995-19508923 A
                            19950313
     DE 1995-19543865 A
                            19951124
     CASREACT 126:100903
QS 
     Novel oligonucleotide analogs which may be loosely described as
AB
     phosphonomonoester analogs of peptide nucleic
     acids (PMENA's) and methods for their synthesis are claimed.
     Particularly preferred PMENA analogs are Q-[OP(:O)(OR)CH2N(COCH2B)CH2CH2]n
     O-Q' (n=1-25; R=OH, OEt, OPh, etc.; B=natural nucleobase; Q,Q'=H, alkyl,
     Ph, etc. or an oligonucleotide or modified oligonucleotide). Their
     application relates to use as inhibitors of gene expression (antisense
     oligonucleotides, ribozymes, sense oligonucleotides and triplex-forming
     oligonucleotides), as probes for the detection of nucleic acids and as
     auxiliaries in mol. biol. PMENA analog H-[OP(:O)(OH)CH2N(COCH2T)CH2CH2]90
     P(:O)(OEt)OEt was prepd. and its interaction with (dA)9 examd. by UV
     spectroscopy and by gel shift anal. The Tm for the PMENA analog-(dA)9
     complex was 23.degree..
ST
     oligonucleotide analog phosphonomonoester synthesis pharmaceutical
     Oligonucleotides
IT
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); PEP
     (Physical, engineering or chemical process); SPN (Synthetic preparation);
     THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study);
     PREP (Preparation); PROC (Process); USES (Uses)
        (analogs; phosphonomonoester nucleic acids prepn. and use in mol. biol.
        and as pharmaceuticals)
     Artery, disease
\operatorname{IT}
        (coronary, restenosis, prevention of; phosphonomonoester nucleic acids
        prepn. and use in mol. biol. and as pharmaceuticals)
    Gene
        (expression, inhibition of; phosphonomonoester nucleic acids prepn. and
        use in mol. biol. and as pharmaceuticals)
     Antitumor agents
IT
     Antiviral agents
        (phosphonomonoester nucleic acids prepn. and use in mol. biol. and as
        pharmaceuticals)
     Probes (nucleic acid)
\operatorname{IT}
     RL: ARG (Analytical reagent use); SPN (Synthetic preparation); ANST
     (Analytical study); PREP (Preparation); USES (Uses)
        (phosphonomonoester nucleic acids prepn. and use in mol. biol. and as
        pharmaceuticals)
IT
     Growth factors, animal
     Tumor necrosis factors
     RL: MSC (Miscellaneous)
        (treatment of diseases involving; phosphonomonoester nucleic acids
        prepn. and use in mol. biol. and as pharmaceuticals)
\operatorname{IT}
     Hepatitis B virus
     Human herpesvirus 1
     Human herpesvirus 2
     Human immunodeficiency virus
     Influenza virus
     Papillomavirus
        (treatment of infection by; phosphonomonoester nucleic acids prepn. and
        use in mol. biol. and as pharmaceuticals)
     185670-74-0P
\operatorname{IT}
     RL: PEP (Physical, engineering or chemical process); SPN (Synthetic
     preparation); PREP (Preparation); PROC (Process)
        (phosphonomonoester nucleic acids prepn. and use in mol. biol. and as
        pharmaceuticals)
IT
     50-00-0, Formaldehyde, reactions 100-27-6 107-18-6, 2-Propen-1-ol,
     reactions 141-43-5, reactions 762-04-9 4712-55-4 14470-28-1
```

20924-05-4 57260-73-8 78635-98-0 89992-70-1 102774-86-7

185670-94-4

172405-18-4 172405-25-3

172405-10-6

```
RL: RCT (Reactant); RACT (Reactant or reagent)
        (phosphonomonoester nucleic acids prepn. and use in mol. biol. and as
        pharmaceuticals)
                  105496-31-9P
     85363-76-4P
                                 183057-32-1P
                                                183057-37-6P
                                                               183057-48-9P
IT
     183057-51-4P 183057-55-8P
                                  183057-59-2P 183057-63-8P
                                                                183057-66-1P
     183057-69-4P 183057-72-9P
                                  183057-75-2P
                                                 183057-79-6P
                                                                183057-82-1P
     183057-84-3P 183057-88-7P
                                  183057-91-2P
                                                 183057-94-5P
                                                                183057-96-7P
     183057-99-0P 183058-02-8P
                                  183058-04-0P
                                                 183058-06-2P
                                                                183058-09-5P
                   183058-11-9P
                                  183058-12-0P
                                                 183058-13-1P
                                                                183058-14-2P
    183058-10-8P
                                  183058-18-6P
                   183058-16-4P
                                                                183058-21-1P
                                                 183058-19-7P
     183058-15-3P
                                  185670-36-4P
     183058-22-2P
                   183058-25-5P
                                                 185670-58-0P
                                                                185670-59-1P
     185670-60-4P
                   185670-61-5P
                                  185670-62-6P
                                                 185670-63-7P
                                                                185670-64-8P
                                                                185670-69-3P
                   185670-66-0P
                                  185670-67-1P
                                                 185670-68-2P
     185670-65-9P
     185670-70-6P
                   185670-71-7P
                                  185670-72-8P
                                                 185670-76-2P
                                                                185670-78-4P
                   185670-80-8P
                                                 185670-82-0P
                                  185670-81-9P
                                                                185670-84-2P
     185670-79-5P
                  185670-90-0P
                                  185670-92-2P
                                                 185670-95-5P
     185670-87-5P
                                                                185670-96-6P
     185670-97-7P 185670-98-8P
                                  185670-99-9P
                                                 185671-00-5P
                                                                185671-01-6P
     185671-02-7P 185671-03-8P
                                  185830-87-9P
                                                 185830-88-0P
                                                                185830-89-1P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (phosphonomonoester nucleic acids prepn. and use in mol. biol. and as
        pharmaceuticals)
    ANSWER 23 OF 26 HCAPLUS COPYRIGHT 2003 ACS
L85
     1996:508642 HCAPLUS
ΑN
       Correction of: 1996:190218
     125:168639
DN
       Correction of: 124:344062
     Synthesis of polyamide nucleic acids (PNAs) using a novel
TI
     Fmoc/Mmt protecting-group combination
     Breipohl, G.; Knolle, J.; Langner, D.; O'Malley, G.;
ΑU
     Uhlmann, E.
     Central Pharma Res., Hoechst AG, Frankfurt, 65926, Germany
CS
     Bioorganic & Medicinal Chemistry Letters (1996), 6(6), 665-670
SO
     CODEN: BMCLE8; ISSN: 0960-894X
PB
     Elsevier
\mathtt{DT}
     Journal
     English
\mathtt{L}\mathtt{A}
CC
     34-3 (Amino Acids, Peptides, and Proteins)
     Section cross-reference(s): 26
AΒ
     The prepn. of 9-fluorenylmethoxycarbonyl (Fmoc) protected building blocks
     for the synthesis of polyamide nucleic acids (PNAs) is
     described. Use of 4-methoxyphenyldiphenylmethyl (Mmt)-protecting groups
     for the exocyclic amino function of the nucleobases enhances the soly. of
     the monomers and allows final deprotection by mild acid treatment. The
     novel synthetic route is exemplified by the synthesis of heptameric and
     octameric PNAs.
    polyamide nucleic acid Merrifield synthesis; peptide
ST
     nucleic acid Merrifield synthesis; monomethoxytrityl
     nucleobase protective group soly
    Merrifield synthesis
ΙT
        (synthesis of peptide nucleic acids using
        a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
        combination)
     Peptide nucleic acids
ΙT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (synthesis of peptide nucleic acids using
        a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
        combination)
     Protective groups
ΙT
        (methoxytrityl, synthesis of peptide nucleic
        acids using a novel fluorenylmethoxycarbonyl and
```

```
monomethoxytrityl protecting group combination)
     71-30-7, Cytosine 73-24-5, Adenine, reactions 96-32-2, Methyl
IT
     bromoacetate 10310-21-1, 2-Amino-6-chloropurine 20924-05-4,
     1-(Carboxymethyl)thymine 172405-43-5
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (synthesis of peptide nucleic acids using
        a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
        combination)
                                  172405-47-9P 172405-48-0P
                                                                172405-49-1P
     169396-92-3P 172405-46-8P
\operatorname{IT}
     172405-50-4P 172405-51-5P 172405-52-6P 172405-53-7P 172405-54-8P
     172405-55-9P 172405-56-0P
                                  172405-57-1P 172405-58-2P 172405-59-3P
     172405-62-8P 176750-53-1P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (synthesis of peptide nucleic acids using
        a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
        combination)
                 172405-67-3P 176750-54-2P
     139166-84-0P
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (synthesis of peptide nucleic acids using
        a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
        combination)
    ANSWER 24 OF 26 HCAPLUS COPYRIGHT 2003 ACS
L85
     1996:190218 HCAPLUS
AN
     124:344062
DN
     Synthesis of polyamide nucleic acids (PNAs) using a novel
{f T}\,{f I}
     Fmoc/Mmt protecting-group combination
     Breipohol, G.; Knolle, J.; Langner, D.; O, Malley, G.;
ΑU
     Uhlmann, E.
    Central Pharma Research, Hoechst AG, Frankfurt, 65926, Germany
     Bioorganic & Medicinal Chemistry Letters (1996), 6(6), 665-70
SO
     CODEN: BMCLE8; ISSN: 0960-894X
     Elsevier
PΒ
     Journal
DT
     English
LA
     34-3 (Amino Acids, Peptides, and Proteins)
CC
     Section cross-reference(s): 26
     The prepn. of 9-fluorenylmethoxycarbonyl (Fmoc) protected building blocks
AΒ
     for the synthesis of polyamide nucleic acids (PNAs) is
     described. Use of 4-methoxyphenyldiphenylmethyl (Mmt)-protecting groups
     for the exocyclic amino function of the nucleobases enhances the soly. of
     the monomers and allows final deprotection by mild acid treatment. The
     novel synthetic route is exemplified by the synthesis of heptameric and
     octameric PNAs.
ST
     polyamide nucleic acid Merrifield synthesis; peptide
     nucleic acid Merrifield synthesis; monomethoxytrityl
     nucleobase protective group soly
     Merrifield synthesis
IT
        (synthesis of peptide nucleic acids using
        a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
        combination)
     Peptide nucleic acids
IT
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (synthesis of peptide nucleic acids using
        a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
        combination)
     Protective groups
IT
        (methoxytrityl, synthesis of peptide nucleic
        acids using a novel fluorenylmethoxycarbonyl and
       monomethoxytrityl protecting group combination)
     71-30-7, Cytosine 73-24-5, Adenine, reactions
                                                     96-32-2, Methyl
IT
```

bromoacetate 10310-21-1, 2-Amino-6-chloropurine 20924-05-4,

```
1-(Carboxymethyl)thymine 172405-43-5
    RL: RCT (Reactant); RACT (Reactant or reagent)
       (synthesis of peptide nucleic acids using
       a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
       combination)
    169396-92-3P 172405-46-8P
                                172405-47-9P 172405-48-0P
                                                            172405-49-1P
    172405-50-4P 172405-51-5P 172405-52-6P 172405-53-7P 172405-54-8P
                                172405-57-1P 172405-58-2P 172405-59-3P
    172405-55-9P 172405-56-0P
    172405-62-8P 176750-53-1P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
       (synthesis of peptide nucleic acids using
       a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
       combination)
    139166-84-0P 172405-67-3P 176750-54-2P
\operatorname{IT}
    RL: SPN (Synthetic preparation); PREP (Preparation)
       (synthesis of peptide nucleic acids using
       a novel fluorenylmethoxycarbonyl and monomethoxytrityl protecting group
       combination)
    ANSWER 25 OF 26 HCAPLUS COPYRIGHT 2003 ACS
L85
AN
    1995:994444 HCAPLUS
    124:202955
DN
    Preparation of polyamide-oligonucleotide derivatives as drugs, gene
TI
    probes, and primers.
    Uhlmann, Eugen; Breipohl, Gerhard
IN
    Hoechst A.-G., Germany
PΑ
    Eur. Pat. Appl., 51 pp.
SO
    CODEN: EPXXDW
\mathsf{DT}
    Patent
LA
    German
    ICM C07H021-00
IC
    ICS C08L077-00; C12Q001-68; A61K031-70
ÇÇ
    33-9 (Carbohydrates)
    Section cross-reference(s): 1, 6, 34
FAN.CNT 1
                                        APPLICATION NO.
    PATENT NO.
                                                        DATE
                    KIND DATE
    EP 672677 A2 19950920
                                        EP 1995-103332
ΡI
                                                       19950308
    EP 672677 A3 19960117
    EP 672677 B1
                          20020703
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
    DE 4408528 A1
                          19950928
                                        DE 1994-4408528 19940314
    EP 1113021 A2 20010704
                                        EP 2001-104012 19950308
    EP 1113021 A3
                          20010711
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE
                                                       19950308
    AT 220070
                          20020715
                                       AT 1995-103332
    ES 2179080
                     T3 20030116
                                        ES 1995-103332
                                                      19950308
    FI 9501132 A 19950915
                                        FI 1995-1132 19950310
    AU 9514798
                    Al 19950921
                                        AU 1995-14798
                                                       19950310
    AU 698210
                     B2 19981029
    CA 2144475
                     AA 19950915
                                        CA 1995-2144475 19950313
    NO 9500955
                     A 19950915
                                        NO 1995-955
                                                       19950313
    CN 1112126
                                                      19950313
                    A 19951122
                                        CN 1995-102946
    JP 07278179 A2 19951024
                                        JP 1995-54644
                                                       19950314
PRAI DE 1994-4408528 A
                         19940314
    EP 1995-103332 A3 19950308
    F[(QB)q(Q1B)r(Q2B)s(Q3B)t]xF1[q, r, s, t = 0, 1; X = 1-20; Q, Q2 = 0]
AΒ
    nucleic acid (deriv.); Q1, Q3 = polyamide residue contg. .gtoreq.1 nucleic
    acid base except thymine; B = covalent bond, org. residue contg. .gtoreq.1
    of C, N, O, S; F, F1 = end groups which may be bound to each other], were
```

prepd. Title compds. show increased cellular uptake, improved nuclease

stability, and are not cytotoxic; they are claimed for use as drugs and

gene probes.

ST polyamide oligonucleotide prepn drug probe primer; dna pna hybrid mol prepn; gene probe polyamide oligonucleotide prepn

IT Neoplasm inhibitors

Nucleic acid hybridization

Virucides and Virustats

(prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

IT Nucleopeptides

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

IT Animal cell

(treatment of diseases influenced by cell-cell adhesion receptors; prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

IT Integrins

RL: BSU (Biological study, unclassified); BIOL (Biological study) (treatment of diseases influenced by integrins; prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

Nucleotides, preparation
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP

(oligo-, prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

IT Nucleotides, preparation

(Preparation); USES (Uses)

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(oligo-, deoxyribo-, prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

IT Heart, disease

(restenosis, treatment; prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

IT 175864-54-7P 175864-55-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BUU (Biological use, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

IT 108-30-5, reactions 502-85-2 4048-33-3, 6-Amino-1-hexanol 20924-05-4 67826-12-4 98796-51-1 100747-20-4 172405-39-9 172405-41-3 172405-42-4 172494-26-7 172494-27-8 172494-28-9

RL: RCT (Reactant); RACT (Reactant or reagent)

(prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

IT 114729-83-8P 125697-62-3P 172316-34-6DP, resin bound 172316-34-6P 172316-40-4P 172316-42-6P 172316-45-9P 172405-31-1P 172494-29-0P 172494-30-3P 172494-31-4P 172494-32-5P 172494-33-6P 172494-34-7P 172494-35-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. of polyamide-oligonucleotide derivs. as drugs, gene probes, and primers)

```
1995:994426 HCAPLUS
AN
    124:87803
DN
    Preparation of substituted N-ethylglycine derivatives for the preparation
TI
    of peptide nucleic acids and peptide
    nucleic acid/deoxyribonucleic acid hybrids.
    Breipohl, Gerhard; Uhlmann, Eugen; Knolle, Jochen
IN
    Hoechst A.-G., Germany
PA
    Eur. Pat. Appl., 31 pp.
SO
    CODEN: EPXXDW
    Patent
\mathtt{DT}
LA
    German
    ICM C07D239-46
IC
    ICS C07D239-54; C07D473-34; C07D473-18; C07D233-92; C07D521-00;
         C08G069-06; C07H021-00; C08G069-10
    34-3 (Amino Acids, Peptides, and Proteins)
CC
    Section cross-reference(s): 33
FAN.CNT 1
    PATENT NO. KIND DATE
                                        APPLICATION NO. DATE
    EP 672661 A1 19950920
                                        EP 1995-103333 19950308
PΙ
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, NL, PT, SE
    DE 4408534 A1 19950928 DE 1994-4408534 19940314
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                                        AU 1995-14799
    AU 9514799 A1 19950921
                                                       19950310
    AU 686729
                     B2 19980212
    CA 2144474
                                        CA 1995-2144474 19950313
                     AA 19950915
    NO 9500959 A
                                        NO 1995-959 19950313
                         19950915
                                        US 1995-402840
    US 6075143 A 20000613
                                                       19950313
    JP 07258222 A2 19951009
                                        JP 1995-54643
                                                       19950314
    US 6465650
                     B1
                                        US 2000-506901
                          20021015
                                                       20000218
PRAI DE 1994-4408534 A
                         19940314
    US 1995-402840
                   А3
                          19950313
    MARPAT 124:87803
OS
    PGXCH2CH2N(COYB)CH2CO2H [PG = urethane- or trityl-type protecting group
AB
    labile to weak acid; X = NH, O, S; Y = CH2, NH, O; B = (protected)
    nucleoside (replacement) base], were prepd. Thus, N-[(4-
    methoxyphenyl)diphenylmethyl]aminoethylglycine Me ester (prepn. given) in
    DMF was treated sequentially with 3,4-dihydro-4-oxo-1,2,3-benzotriazine,
    4-ethylmorpholine, N4-benzoyl-N1-carboxymethylcytosine in DMF, and with
    DCC; the mixt. was stirred 20 h at room temp. to give the coupling
    product, which was sapond. with aq. NaOH/dioxane to give
    N-[(4-methoxyphenyl)diphenylmethyl]aminoethyl-N-[[1-(N4-
    benzoyl)cytosyl]acetyl]glycine.
    ethylglycine deriv pna intermediate prepn; dna pna
ST
    hybrid intermediate ethylglycine deriv; nucleopeptide intermediate
    ethylglycine prepn
    Deoxyribonucleic acids
IT
    RL: SPN (Synthetic preparation); PREP (Preparation)
       (hybrids; prepn. of substituted N-ethylglycine derivs. for the prepn.
       of peptide nucleic acids and
       peptide nucleic acid/DNA hybrids)
IT
    Nucleopeptides
    RL: SPN (Synthetic preparation); PREP (Preparation)
       (intermediates; prepn. of substituted N-ethylglycine derivs. for the
       prepn. of peptide nucleic acids and
       peptide nucleic acid/DNA hybrids)
    65-71-4, Thymine 71-30-7, Cytosine 73-24-5, Adenine, reactions
ΙT
    73-40-5, Guanine 79-04-9 79-08-3, Bromoacetic acid 96-32-2
    98-88-4, Benzoyl chloride 100-07-2, 4-Methoxybenzoyl chloride
    141-43-5, 2-Aminoethanol, reactions 156-57-0, 2-Mercaptoethylamine
    hydrochloride 288-88-0, 1H-1,2,4-Triazole 298-12-4, Glyoxylic acid
    794-94-5, 4-Methoxybenzoic anhydride 1710-98-1 3034-38-6,
    4-Nitroimidazole 3587-60-8, Benzyl chloromethyl ether 18907-79-4
```

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40615-36-9 67826-12-4 112233-74-6
    34619-03-9, Di-tert-butylcarbonate
    RL: RCT (Reactant); RACT (Reactant or reagent)
        (prepn. of substituted N-ethylglycine derivs. for the prepn. of
       peptide nucleic acids and peptide
       nucleic acid/DNA hybrids)
    79-30-1P, Isobutanoyl chloride 13251-16-6P 20924-05-4P
                                                                21047-89-2P
ΙT
    26661-13-2P 51820-70-3P 55036-34-5P 97025-97-3P 118534-11-5P
    119451-90-0P 134456-94-3P 135697-25-5P 141743-19-3P
                                                              168263-86-3P
    170944-06-6P 172405-08-2P 172405-09-3P 172405-10-6P
                                                              172405-11-7P
    172405-12-8P 172405-13-9P 172405-14-0P 172405-15-1P
                                                              172405-16-2P
                                 172405-19-5P 172405-20-8P
    172405-17-3P 172405-18-4P
                                                              172405-21-9P
    172405-22-0P 172405-23-1P 172405-24-2P 172405-25-3P
                                                              172405-26-4P
    172405-27-5P 172405-28-6P 172405-29-7P 172405-30-0P
                                                               172405-38-8P
                                 172405-41-3P 172405-42-4P
    172405-39-9P 172405-40-2P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (prepn. of substituted N-ethylglycine derivs. for the prepn. of
       peptide nucleic acids and peptide
       nucleic acid/DNA hybrids)
    170490-73-0P 172316-36-8P 172316-40-4P 172316-41-5P
                                                              172316-42-6P
IT
    172316-44-8P 172316-45-9P 172405-31-1P 172405-32-2P
                                                               172405-33-3P
    172405-34-4P 172405-35-5P 172405-36-6P 172405-37-7P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (prepn. of substituted N-ethylglycine derivs. for the prepn. of
       peptide nucleic acids and peptide
       nucleic acid/DNA hybrids)
=> d his
     (FILE 'HOME' ENTERED AT 07:36:09 ON 13 MAR 2003)
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             1 S E3, E4
L1
               SEL RN
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            88 S E1-E88
L2
             0 S L2 AND (NCNC2-SC4 AND NCNC2-NCNC3 AND NCNC3)/ES
L3
             0 S L2 AND NCNC2-SC4/ES
L4
L5
             6 S L2 AND P/ELS
            86 S L2 AND SQL/FA
L6
            17 S L6 AND 11/SQL
Ļ7
\Gamma8
            26 S L6 AND 12/SQL
L9
             4 S L8 AND PEPTIDE NUCLEIC ACID AND THIENO AND IMIDAZOL AND HEXAH
L10
             1 S L9 AND G G T A T G G G A T A T
               E FS
               E GGTATGGGATAT/SQEN
             3 S E3
L11
               E TATTCCGTCAT/SQEN
           129 S E3
L12
L13
             4 S L12 AND THIENO AND IMIDAZOL?
L14
             2 S L13 NOT 22/SQL
               E TATTCCGTCAT/SQEN
             2 S L2 NOT L6
L15
L16
             7 S L2 AND ?THIEN?/CNS
             4 S L2 AND ?GUAN?/CNS
L17
             1 S L2 AND ?ADEN?/CNS NOT L17
L18
             1 S L2 AND ?THYM?/CNS NOT L17, L18
L19
            41 S (NCNC2-SC4 AND NCNC2-NCNC3 AND NCNC3)/ES
L20
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L21

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0 S L21 AND 1/P
L22
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L23
L24
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L25
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L26
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L27
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L28
             127 S L28 AND OXOPENTYL AMINO HEXYL
L29
L30
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L31
               4 S L30 NOT L31
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               0 S L32 NOT OC4/ES
L33
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L35
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L36
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· L39
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L40
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L42
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L44
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L50
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L51
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               O S L52 AND NCNC2-SC4/ES
L53
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L54
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L55
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L56
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L57
              2 S L54 AND ACETYL
L58
              12 S L11, L14, L40, L44, L48, L49, L56, L58
L59
                 SAV L59 SIEW835A/A
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L60
                 E UHLMANN E/AU
             173 S E3, E4, E14-E15
L61
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                 E BREIPOHL G/AU
             106 S E3-E6
L62
                 E BREIPOEHL G/AU
               1 S E2
L63
L64
               1 S E10
                 E WILL D/AU
              40 S E3, E7-E10
L65
                 E AVENTIS/PA, CS
            1598 S E2-E4
L66
             857 S (AVENTIS(L) PHARM?)/PA,CS
L67
L68
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               3 S L60, L68
L69
                 E PEPTIDE NUCLEIC ACID/CT
                 E E4+ALL
            1670 S E3
L70
            5997 S PEPTIDE NUCLEIC ACID OR PNA
L71
              34 S L61-L67 AND L70,L71
L72
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SEL RN L72

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L74 0 S L73 AND NCNC2-SC4/ES				
L75 8 S L73 AND (?THIENO?(L)?IMIDAZ	Z?)/CNS			
L76 27 S L73 AND L11, L12				
L77 7 S L73 AND L52				
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L79 9 S L78 AND L61-L67				
L80 8 S L79 AND L72				
L81 9 S L79, L80				
L82 3 S L78 NOT L81				
FILE 'REGISTRY' ENTERED AT 08:50:05 ON 1	13 MAR 2003			
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SEL HIT RN L69	'HCAPLUS' ENTERED AT 08:50:26 ON 13 MAR 2003 SEL HIT RN L69			
DIT TO INTOTOTION TO THE AND A COLUMN TO THE AND A	10 MAD 0000			
FILE 'REGISTRY' ENTERED AT 08:50:55 ON 13 MAR 183 11 S E565-E575				
				FILE 'HCAPLUS' ENTERED AT 08:51:21 ON 13 MAR 2
L84 12 S L78-L82	7 11111 2000			
L85 26 S L72 NOT L69, L84	•			